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Utilizing Targeted Gene Therapy with Nanoparticles Binding Alpha v Beta 3 for Imaging and Treating Choroidal Neovascularization

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Abstract

Purpose: The integrin αvβ3 is differentially expressed on neovascular endothelial cells. We investigated whether a novel intravenously injectable αvβ3 integrin-ligand coupled nanoparticle (NP) can target choroidal neovascular membranes (CNV) for imaging and targeted gene therapy.

Methods: CNV lesions were induced in rats using laser photocoagulation. The utility of NP for in vivo imaging and gene delivery was evaluated by coupling the NP with a green fluorescing protein plasmid (NP-GFPg). Rhodamine labeling (Rd-NP) was used to localize NP in choroidal flatmounts. Rd-NP-GFPg particles were injected intravenously on weeks 1, 2, or 3. In the treatment arm, rats received NP containing a dominant negative Raf mutant gene (NP-ATPμ-Raf) on days 1, 3, and 5. The change in CNV size and leakage, and TUNEL positive cells were quantified.

Results: GFP plasmid expression was seen in vivo up to 3 days after injection of Rd-NP-GFPg. Choroidal flatmounts confirmed the localization of the NP and the expression of GFP plasmid in the CNV. Treating the CNV with NP-ATPμ-Raf decreased the CNV size by 42% (P < 0.001). OCT analysis revealed that the reduction of CNV size started on day 5 and reached statistical significance by day 7. Fluorescein angiography grading showed significantly less leakage in the treated CNV (P < 0.001). There were significantly more apoptotic (TUNEL-positive) nuclei in the treated CNV.

Conclusion: Systemic administration of αvβ3 targeted NP can be used to label the abnormal blood vessels of CNV for imaging. Targeted gene delivery with NP-ATPμ-Raf leads to a reduction in size and leakage of the CNV by induction of apoptosis in the CNV.

Introduction

Age-related macular degeneration (AMD) is the leading cause of blindness in developed countries for people over the age of 50 [1–3]. The neovascular or “wet” form of the disease, characterized by the development of choroidal neovascular membranes (CNV) is the main cause of visual impairment in macular degeneration [3–5]. With the advent of new treatment options such as photodynamic therapy, and especially intravitreal antiangiogenic pharmacotherapy, the visual prognosis of patients with CNV has improved significantly [6–9]. However, the current standard-of-care therapies require monthly intravitreal injections by a retina specialist due to their short half-life in the vitreous [10,11]. Aside from the logistic difficulties and the patients’ discomfort, it also puts the patient at risk for cataract formation, endophthalmitis, vitreous hemorrhage, and retinal detachment. Thus, there is a great need for alternative means of delivering antineovascular therapy to the retina.

Recently, there has been substantial progress in the development of nanoparticles with an integrin-targeted delivery system [12–15]. During vascular remodeling and angiogenesis, several integrins are expressed on the endothelial cells to potentiate cell invasion and proliferation [16,17]. Among them, integrin αvβ3 is expressed on many cell types but its expression level in normal tissue is generally low [18,19]. It is preferentially expressed on angiogenic blood vessels, mediating survival signal and facilitating vascular cell proliferation [20,21]. Previous reports show that integrin αvβ3 is involved in ocular angiogenesis [22,23]. In vivo experiments have shown antibodies blocking or immunoconjugate drug therapy targeting integrin αvβ3 inhibit neovascularization.
In addition, integrin αvβ3 potentiates the internalization of various viruses [27,28], making it a potential target for drug delivery via liposome based nanoparticles.

Previously we have shown that systemic injection of a cationic nanoparticle coupled to an integrin αvβ3-targeting ligand (NP) can deliver a suicide gene to the tumor neovasculature in rats, causing apoptosis and significant tumor regression [12]. Here we evaluated and were able to demonstrate that NP can target choroidal neovascular membranes (CNV) in rats for imaging and targeted gene therapy using a plasmid DNA encoding ATPm-Raf, a dominant-negative mutant form of Raf kinase [29].

Materials and Methods

Animals and Ethics Statement

All experiments were conducted in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health, and the guidelines established by the Animal Care Committee (ACC) of the Massachusetts Eye and Ear Infirmary. The protocol was approved by the ACC (protocol number 07-10-012). A total of 106 Brown-Norway male rats weighing 175–225 grams were obtained from Charles River Laboratories (Wilmington, MA) and used for the experiments.

Characteristics and preparation of Nanoparticles

Detailed description of the NPs and their synthesis has been published previously [12]. All custom-made lipids and genes were GLP manufactured. Briefly, purified lipid components were dissolved in organic solvents (CHCl3 and CH3OH in a ratio 1:1). The CHCl3 and CH3OH were evaporated and dried in rotavap for 24 hours. Distilled and deionized water was added to yield a heterogeneous solution of 30 mM in total lipid concentration. The lipid/water mixture was then sonicated with a probe-tip sonicator for at least one hour. Throughout sonication, the pH of the solution was maintained between 7.0 and 7.5 with 0.01N NaOH solution, and the temperature was maintained above the gel-liquid crystal phase transition point (Tm). The liposome solution was transferred to a petri dish resting on a bed of wet ice, cooled to 0°C, and irradiated at 254 nm for at least one hour with a hand-held UV lamp placed 1 cm above the petri dish, yielding NPs. The NPs were then filtered through a 0.2 µm filter and collected.

Using a Brookhaven dynamic light scattering system (DLS), the size (diameter), distribution, and zeta potential of NPs were determined to be 45.3±2.4 nm and +35mV respectively, averaged for 17 cycles of NP synthesis.

The rats received intravenous treatments at a dose 1 mg/kg of NP and 1 µg/kg of plasmid DNA containing the Raf mutant gene (ATPm-Raf). Total volume of injection was 350 µl.

Induction of Choroidal Neovascular Membranes

Animals were anesthetized with an intraperitoneal injection of 0.2 to 0.3 mL of a 1:1 mixture of 100 mg/mL ketamine and 20 mg/mL xylazine. Pupils were dilated with 5.0% phenylephrine and 0.8% tropicamide. CNV was induced in the eyes of rats with a 532-nm laser (Oculight GLx; Iridex, Mountain View, CA), as previously described [30–32]. Four to eight laser spots (180 mW, 100 µm, 100 ms) were placed in each eye of the rat with a slit lamp delivery system and a cover slip serving as a contact lens. If significant hemorrhage occurred, the eye was excluded.

Evaluation of Specific Targeting of CNV Using In Vivo Imaging and Choroidal Flatmounts

NP carrying a green fluorescing protein (GFP) plasmid (NP-GFP) was used to evaluate the ability of the particles to deliver a gene to the neovascular endothelial cells. Rhodamine labeling of the NP (Rd-NP) was used to localize the particles in choroidal flatmounts. Rats were divided into 3 groups of 6 animals and evaluated 1, 2, or 3 weeks after creation of CNV. The formation of CNV was maintained between 7.0 and 7.5 with 0.01N NaOH solution, and the temperature was maintained above the gel-liquid crystal phase transition point (Tm). The liposome solution was transferred to a petri dish resting on a bed of wet ice, cooled to 0°C, and irradiated at 254 nm for at least one hour with a hand-held UV lamp placed 1 cm above the petri dish, yielding NPs. The NPs were then filtered through a 0.2 µm filter and collected.

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Figure 1. Bioimaging with NP-angiography showing GFP expression using the Topcon camera with fluorescein angiography filter settings. Late phase FAs (A and D) show the CNV lesions prior to injection of NP. Autofluorescent images taken prior to injection of NP reveal minimal background fluorescence of the CNV lesions (B and E). Injection of targeted NP carrying a GFP plasmid (NP-GFP) causes increased fluorescence of the CNV lesions from GFP expression (C) whereas non-targeted NP carrying a GFP plasmid (ntNP-GFP) does not cause any increase in the intensity of fluorescence of the CNV over background autofluorescence (F).

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CNV was confirmed by fluorescein angiography (FA) using a digital fundus camera (TCC501A; Topcon, Paramus, NJ) following intraperitoneal injection of 0.2 mL of 2% fluorescein sodium. The rats were then injected with in vivo imaging using the same camera and FA filter settings 24, 48, and 72 hours after injection of particles. The rats were then euthanized and choroidal flat mounts were performed at the above time points. RPE-choroid-sclera complex was flatmounted (Vector Laboratories, Burlingame, CA) and coverslipped. Pictures of the choroidal flatmounts were taken by a confocal microscope (Leica Microsystems, Wetzlar, Germany). Negative controls were evaluated under identical conditions without injection of NP or following injection of non-targeted rhodamine labeled liposome particles carrying GFP-plasmid.

**Treatment of CNV with Targeted Gene Delivery**

After creation of laser-induced CNV, animals were divided to 7 groups; groups A–C were treatment groups and groups D–G were controls. Group A (n = 6) received one intravenous injection of \( \alpha \beta \) targeted-NP containing ATP\( \mu \)-Raf (NP-ATP\( \mu \)-Raf) on days 1, 3, and 5 after CNV creation; group B (n = 6) received one intravenous injection of NP-ATP\( \mu \)-Raf on days 3, 5, and 7; group C (n = 3) received one intravenous injection of NP-ATP\( \mu \)-Raf on days 7, 9, and 11; group D (n = 6) did not receive any treatment; group E (n = 3) received one intravenous injection of non-targeted NP containing ATP\( \mu \)-Raf (ntNP-ATP\( \mu \)-Raf) on days 1, 3, and 5; group F (n = 3) received one intravenous injection of \( \alpha \beta \) targeted-NP without ATP\( \mu \)-Raf (NP) on days 1, 3, and 5; and group G (n = 3) received one intravenous injection of ATP\( \mu \)-Raf gene without NP on days 1, 3, and 5.

**Evaluation of CNV Size and Leakage**

We used FA and OCT to monitor CNV development and changes in vivo and choroidal flatmounts to study the size of the lesions ex vivo. FA was performed as detailed above, 1 and 2 weeks after CNV creation in all treated and control animals. A choroidal neovascular membrane was defined as fully regressed after treatment if there was no leakage in the area of treated membrane [30,31]. The angiograms were graded by two masked readers using a pre-established grading scheme [33,34]. Briefly, the description of each grade follows: 0, faint hyperfluorescence or mottled fluorescence without leakage; 1, hyperfluorescent lesion without progressive increase in size or intensity; 2A, hyperfluorescence increasing in intensity but not in size; 2B, hyperfluorescence increasing in intensity and in size.

Two weeks after CNV creation, the size of the CNV lesions was measured in choroidal flatmounts using the methods reported previously after perfusion of 5 mg/mL fluorescein labeled dextran [31,35]. A computer program (OpenLab; Improvision, Boston, MA) was used by two masked investigators to measure the hyperfluorescent areas corresponding to the CNV lesions.

**Optical Coherence Tomography Measurement of CNV Size**

Six rats treated with intravenous NP-ATP\( \mu \)-Raf on days 1,3 and 5 and 6 control rats without treatment were used for evaluation of CNV size in vivo using optical coherence tomography (SDOCT, Bioptigen, Durham, NC) on days 3, 5, and 7 after CNV creation. A volume analysis was performed, using 100 horizontal raster, consecutive B-scan lines, each one composed of 1200 A-scans. The volume size was 2.1 x 2.1 cm. To evaluate the cross-sectional size of each lesion in OCT images, the sections passing through the center of the CNV were chosen. The center of the lesion was defined as the midline passing through the area of RPE-Bruch’s membrane rupture. In order to consistently identify this point, we used the en-face fundus reconstruction tool provided with the Bioptigen SD-OCT system. For each time point, the same spot was used to evaluate the size of the CNV. CNV was outlined from the inner border of the retinal pigment epithelial layer to the top of the lesion and the size was measured using Image J software (http://rsbweb.nih.gov/ij/, last access January 7th 2009).

**Histopathology of CNV Lesions**

On day 3, 5, 7 and 14 after CNV creation, eyes were enucleated and fixed in 4% paraformaldehyde in phosphate-buffered saline.
(PBS) for 1 hour and cryoprotected. Serial sections of the eyes were cut at 10 μm thickness on a cryostat (CM1850; Leica, Heidelberg, Nussloch, Germany) at −20°C, and prepared for staining. Terminal deoxyuridine triphosphate Nick-End Labeling (TUNEL) assay was performed according to the manufacturer's protocol (ApoTag Fluorescein in situ Apoptosis Detection Kit; Chemicon, Temecula, CA) as previously reported [31,36]. CD31 (1:100, Serotec, Oxford, UK) antibody was used for visualizing endothelial cells and a mouse monoclonal antibody for ED1, the rat homologue of human CD68 (1:100, Millipore, Billerica, MA) was used for staining macrophages. Sections were then stained with DAPI (1:1000, Invitrogen Ltd, Carlsbad, CA, USA) for nuclear staining and mounted with Vecta shield mounting media (Vector Laboratories, Burlingame, CA). Photographs of the CNV were taken with upright fluorescent microscope (DM RXA; Leica, Solms, Germany) and the number of TUNEL positive and ED 1 positive cells were counted.

**Statistical Analysis**

All values are presented as mean ± SE. Paired groups were compared using the Wilcoxon t-test. For three groups, data were compared by Kruskal-Wallis test and for two group comparisons,

![Figure 3. Late phase fluorescein angiography (FA) and choroidal flatmounts (x10) two weeks after laser photocoagulation.](image)

Representative lesions are from the control group (A–D) and the NP-ATPα-Raf treated group (E and F). Group (A) received no treatment; (B) received intravenous injection of non-targeted NP containing ATPα-Raf on days 1, 3, and 5 after laser CNV creation; (C) received intravenous injection of α,β3 targeted NP without ATPα-Raf gene on days 1, 3, and 5; (D) received injection of ATPα-Raf gene without NP on days 1, 3, and 5; (E) received injection of α,β3 targeted-NP containing ATPα-Raf (NP-ATPα-Raf) on days 1, 3, and 5; and (F) received injection of NP-ATPα-Raf on days 3, 5, and 7. NP-ATPα-Raf treated groups (E and F) had significantly lower grade CNV lesions on FA grading and smaller CNV size compared to the control group (A–D). No statistically significant difference in size was noted between the control groups A–D. Quantification of the CNV size on choroidal flat mounts is shown in (G). *P<0.01. Data are expressed as the mean ± SE.

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Mann-Whitney U test was used (SPSS statistics 17.0, SPSS Inc., Chicago, IL, USA). A P value of less than 0.05 was considered statistically significant.

**Results**

**In Vivo imaging Reveals GFP-Plasmid Expression by CNV**

Formation of CNV was confirmed by FA prior to injection of NP. One day after injection of rhodamine-labeled αβ3 targeted nanoparticle carrying a GFP plasmid (Rd-NP-GFPg), digital fundus photography with FA filter settings revealed hyperfluorescence of the CNV lesions, confirming localization and adhesion of the NP and GFP expression. This hyperfluorescence was sustained through day 3 (data not shown) and was seen in 1, 2, and 3-week-old CNV lesions examined (Figure 1A–C). There was no notable hyperfluorescence noted above background fluorescence level in the control groups (Figure 1D–F). There was no evidence of increased fluorescence in the normal retinal or choroidal vasculature.

**Rhodamine-Labeled αβ3 Targeted Nanoparticle Accumulate in the CNV and Induce GFP Expression**

The delivery of rhodamine dye and the expression of GFP plasmid in the CNV were confirmed by performing confocal microscopy on choroidal flat mounts (Figure 2). Choroidal flatmounts performed after in vivo imaging revealed accumulation of rhodamine labeled nanoparticles in the CNV lesion (Figure 2C). There was an overlap of GFP expression and the rhodamine accumulation in the CNV (Figure 2D). No notable difference was seen in the pattern of NP accumulation or intensity of GFP expression over the time course examined. There was no evidence of increased fluorescence in the normal choroidal vasculature and no evidence of rhodamine accumulation or GFP expression in the control animal injected with non-targeted rhodamine labeled nanoparticles carrying a GFP plasmid (Rd-ntNP-GFPg; Figure 2E–H).

**αβ3 Targeted-NP Containing ATPμ-Raf Reduces the Size of CNV**

Treatment of CNV with intravenous injection of αβ3 targeted-NP containing ATPμ-Raf (NP-ATPμ-Raf) on days 1, 3, and 5 resulted in a 42.0% reduction in the CNV size on choroidal flatmount compared with control CNVs with no treatment (mean size, 53338.7 μm² vs. 31029.3 μm², p<0.001) (Figure 3A and E).

![Figure 4. In vivo evaluation of CNV utilizing SD-OCT.](image-url)

**Reduced Endothelial Cell Count and Increased Apoptosis in CNV Treated with NP-ATPμ-Raf**

CD31-positive endothelial cells were detected in the subretinal space starting on day 3 after laser injury (figure 5D). The cells increased and focalized to a distinct subretinal membrane representing the CNV by day 7. In the treated CNV, there were

![Figure 4. In vivo evaluation of CNV utilizing SD-OCT.](image-url)
fewer CD31-positive cells and the CNV appeared more compact with distinct borders and fibrous formation by day 7 (Figure 5D).

We investigated signs of apoptosis with TUNEL staining. In the treated group, significantly more TUNEL-positive nuclei were observed in the CNV starting on day 3 (Figure 5A and B). This trend continued through day 7 (Mann Whitney U test, p<0.01 for day 3 and 5, p=0.01 for day 7, Figure 6B). The reduction of the CNV size as measured in histological sections reached statistical significance on day 7 (Mann Whitney U test, P=0.001, Figure 6C).

Macrophage infiltration with ανβ3 Targeted-NP Containing ATPμ-Raf
ED1 positive cells (a marker for macrophages, equivalent to human CD68) were concentrated within the subretinal space at the laser injury site on day 3 (Figure 6D). No ED1-positive cells were observed in the undamaged choroid. No difference was noted in the number of ED1-positive cells infiltrated into the CNV between the treated group and the untreated group on day 3. However, on day 5 and 7 statistically more ED1-positive cells were seen in the treatment group (Mann Whitney U test, P<0.01 respectively; Figure 6D).

Discussion
Specific targeting and delivery of medication for the treatment of CNV remains challenging [37]. In this study we were able to target experimental CNV after systemic injection of a cationic

Table 1. Fluorescein angiography (FA) grading of CNV lesions.

<table>
<thead>
<tr>
<th>Grade 1</th>
<th>Grade 2A</th>
<th>Grade 2B</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>5(6.2%)</td>
<td>16(19.8%)</td>
<td>60(74.1%)</td>
</tr>
<tr>
<td>mNP-ATPμ-Raf</td>
<td>2(5%)</td>
<td>10(25%)</td>
<td>28(70%)</td>
</tr>
<tr>
<td>ATPμ-Raf</td>
<td>3(7.1%)</td>
<td>7(16.7%)</td>
<td>32(76.2%)</td>
</tr>
<tr>
<td>NP</td>
<td>0(0%)</td>
<td>6(16.2%)</td>
<td>31(83.8%)</td>
</tr>
<tr>
<td>NP-ATPμ-Raf on D1, 3, 5</td>
<td>10(13.9%)</td>
<td>33(45.8%)</td>
<td>59(40.3%)</td>
</tr>
<tr>
<td>NP-ATPμ-Raf on D3, 5, 7</td>
<td>1(2.8%)</td>
<td>16(44.4%)</td>
<td>19(52.8%)</td>
</tr>
</tbody>
</table>

Significantly higher percentage of control CNV lesions were Grade 2B compared to the treated CNV (Pearson Chi-square test P<0.001). There were no significant differences between the different treatment groups (Pearson Chi-square test P=0.679).

doi:10.1371/journal.pone.0018864.t001

Figure 5. Evaluation of endothelial cell apoptosis with TUNEL staining in frozen sections. Quantification of TUNEL positive cells showed significantly more TUNEL(+) cells/lesion (A) and TUNEL (+) cells/mm² (B) with treatment of NP-ATPμ-Raf compared to the control group on day 3 and 5 after laser injury. There was a statistically significant reduction of CNV size noted on day 7(C). Double-immunofluorescent staining of frozen sections (x20) obtained at 3, 5 and 7 days after laser photocoagulation for the endothelial cell marker CD31 and TUNEL stain (D). *P<0.01. Data are expressed as the mean ± SE.
doi:10.1371/journal.pone.0018864.g005
A nanoparticle coupled to an integrin $\alpha_v\beta_3$-targeting ligand (NP) and utilize this method for imaging and treatment of CNV in rats.

We first demonstrated the vascular targeting of NP and its ability to deliver a gene to the neovascular endothelial cells of CNV in rats using rhodamine labeled NPs coupled with GFP (Rd-NP-GFPg). GFP expression was seen in vivo using fundus NP-angiography with FA filter settings and ex vivo in choroidal flat mounts. In vivo imaging revealed increased fluorescence, sustained for 3 days after NP injection, in 1, 2, or 3 week old CNV (figure 1). GFP expression co-localized to the area of rhodamine labeled NP accumulation in the CNV on choroidal flat mounts. In vivo imaging revealed increased fluorescence, sustained for 3 days after NP injection, in 1, 2, or 3 week old CNV (figure 1). The specific targeting of NP is due to the selectivity of its binding ligand for integrin $\alpha_v\beta_3$ [12], which has limited cellular distribution in normal tissue including the eye [23,38]. This integrin is significantly up regulated during the process of vascular remodeling and angiogenesis and is present in pathologic specimens from human eyes with CNV or proliferative diabetic retinopathy (PDR) [17,22,23,39,40]. With the specific labeling of CNV demonstrated here, fundus NP-angiography utilizing various dyes has the potential to be a novel imaging technique for detecting new CNV or to follow CNV activity independent of CNV size and leakage. Recently, Takeda and colleagues have shown early detection of experimental CNV, not visible on FA, using a similar technique with anti-CCR3 antibody fragments [41]. The advantage of using NPs over immunoconjugate dyes in bioimaging and targeted therapy is that they are less likely to incite an immune reaction.

In the treatment arm of the study, we were able to demonstrate significant reduction of CNV size and leakage by targeted gene therapy using NP coupled to ATP$_\mu$-Raf, a dominant negative form of Raf kinase (Figure 3, 4). Previously Singh et al treated experimental CNV using poly-lactide-co-glycolide (PLGA) nanoparticles carrying a VEGF inhibitory gene [42]. Our approach is different in a number of ways including the type of particles used, specificity of targeting, and the gene delivered. The particles that Singh et al used had a negative z-potential as expected for PLGA nanoparticles, our therapy has a neutral charge and therefore is not toxic in tissue-cultured cells. Our nanoparticles are smaller (45.3 nm vs. 270–420 nm) and form a stable shell through covalent bonds that are far more stable in blood circulation than polymer based (used by Singh et al and hydrolyzed in aqueous solutions).

Figure 6. Increased macrophage infiltration at the site of treated CNV. Macrophage infiltration was highest on day 3 with gradual decrease on days 5 and 7. Significantly higher number of macrophages were observed with the NP-ATP$_\mu$-Raf treated group compared to the control group on days 5 and 7 (A and B). There was a statistically significant reduction of CNV size noted on day 7(C). Immunofluorescent staining of representative frozen sections ($\times$20) obtained at 3, 5, and 7 days after laser photocoagulation for ED 1, a marker for macrophage (D). *$P<0.01$. Data are expressed as the mean $\pm$ SE. doi:10.1371/journal.pone.0018864.g006
environment) or liposome based delivery systems. This stability may lead to more efficacy and/or more side effects. A separate toxicity study is planned to answer this question. The targeted gene, Raf kinase, is an integral member of an intracellular signal transduction pathway involved in regulation of the cell cycle. ATP1-Raf is a mutant form of Raf-1 that fails to bind ATP and blocks the endothelial cell Raf activity in vitro [12,29]. Raf-1 mutation has been linked to vascular defect and apoptosis during embryogenesis and gene therapy with ATP1-Raf causes endothelial cell apoptosis and tumor regression in rats [12,43]. Our results indicate a similar mechanism of CNV regression, through induction of apoptosis in neovascular endothelial cells, after targeted gene therapy with NP-ATP1-Raf. There were significantly higher number of TUNEL-positive cells in the treated CNV as compared to the controls (Figure 5). Moreover, while the recruitment of macrophages per lesion decreased with time in both groups, more macrophage infiltration was noted on days 5 and 7 in the treatment group (Figure 6). The initial spike in macrophage infiltration is likely the result of the inflammatory response to the laser injury. The increased macrophage recruitment to the treated CNV closely follows the increased apoptotic activity, suggesting that the macrophages may be responding to cell death by apoptosis or necrosis.

Repeated systemic administration of NP for the treatment of CNV may lead to side effects such as blood clots in the elderly patient population. However, Intravenous (I.V.) administration of our treatment is not repetitious as opposed to the intravitreal injections of anti-VEGF therapies for example. In addition, for some patients, I.V. injections are less invasive than intravitreal injections and do not carry the potential ocular complications. Furthermore, due to the specificity of the binding site, the dose of nanoparticles needed is small and the tissue distribution limited, making systemic side effects less likely. In our study, we did not attempt the intravitreal route of delivery as part of the investigation. Due to the large size of the NP (45.3 nm), it is unclear if it can distribute through the vitreous and cross the retina to reach the lumen of the CNV vessels through phagocytosis, pinocytosis or other mechanisms.

We also explored the timing of imaging and treatment with targeted NP in our study. Although we were able to target CNV with NP and capture GFP expression by NP-angiography up to three weeks after CNV induction, the maximum efficacy of treatment was achieved when treatment was given on days 1,3 and 5 (42% reduction in CNV size). We saw a modest decrease in the CNV size (24.6%) when the treatment was given on days 3, 5 and 7 and found no effect with treatment on days 7, 9 and 11. This data is consistent with the timing of integrin αvβ3 expression in the laser induced CNV in rats [44]. Integrin αvβ3 is expressed up to 4 weeks after laser injury, however its expression levels are significantly higher in the early stages of CNV formation, peaking at day 7 [44]. The fact that we were unable to show CNV regression with treatment starting at later time points may be due to the concentrations of NP used or the efficacy of transfecting the CNV endothelial cells with ATP1-Raf. Integrin αvβ3 expression has been shown in pathologic specimens from human CNV and diabetic retinopathy [23,45], however, the timing and duration of expression of this integrin has not been studied.

In summary, our results provide evidence that systemic administration of αvβ3 targeted NP can be used to label the abnormal blood vessels of CNV for imaging and targeted gene therapy with ATP1-Raf. These results provide a proof-of-concept for this emerging technology and encourage further experimentation to discern the integration of NPs into current imaging and treatment of CNV and retinal neovascularization. Large animal studies and experiments utilizing NP coupled to various dyes and chemotherapeutic or anti-vascular compounds need to be pursued to further explore the efficacy of this diagnostic and treatment strategy.

Author Contributions
Conceived and designed the experiments: HS-H SG JWM TH DV. Performed the experiments: HS-H MIR AG TH SN IKK ESG SG. Analyzed the data: HS-H MIR AG SG JWM DV. Contributed reagents/materials/analysis tools: SG JWM DV TH SN. Wrote the paper: HS-H MIR AG SG JWM DV.

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