Thermal Stimulation of the Retina Reduces Bruch’s Membrane Thickness in Age Related Macular Degeneration Mouse Models

Jan Tode1,*, Elisabeth Richert1,*, Stefan Koinzer1, Alexa Klettner1, Claus von der Burchard1, Ralf Brinkmann2, Ralph Lucius3, and Johann Roider1

1 Christian-Albrechts-University of Kiel, University Medical Center, Department of Ophthalmology, Kiel, Germany
2 Institute for Biomedical Optics, University of Lübeck, and Medical Laser Center Lübeck GmbH, Lübeck, Germany
3 Christian-Albrechts-University of Kiel, Institute of Anatomy, Kiel, Germany

Correspondence: Jan Tode, Christian-Albrechts-University of Kiel, University Medical Center, Department of Ophthalmology, Arnold-Heller-St 3, 24105 Kiel, Germany. e-mail: jan.tode@uksh.de

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Purpose: To investigate the effect of thermal stimulation of the retina (TS-R) on Bruch’s membrane (BrM) thickness in age-related macular degeneration (AMD) mouse models as a novel concept for the prophylaxis and treatment of dry AMD.

Methods: Two knockout AMD mouse models, B6.129P2-Apoetm1Unc/J (ApoE−/−) and B6.129X1-Nfe2I2tm1Ywk/J (NRF2−/−), were chosen. One randomized eye of each mouse in four different groups (two of different age, two of different genotype) of five mice was treated by TS-R (532 nm, 10-ms duration, 50-μm spot size), the fellow eye served as control. Laser power was titrated to barely visible laser burns, then reduced by 70% to guarantee for thermal elevation without damage to the neuroretina, then applied uniformly to the murine retina. Fundus, optical coherence tomography (OCT), and fluorescein angiography (FLA) images were obtained at the day of treatment and 1 month after treatment. Eyes were enucleated thereafter to analyze BrM thickness by transmission electron microscopy (TEM) in a standardized blinded manner.

Results: Fundus images revealed that all ApoE−/− and NRF2−/− mice had AMD associated retinal alterations. BrM thickness was increased in untreated controls of both mouse models. Subvisible TS-R laser spots were not detectable by fundus imaging, OCT, or FLA 2 hours or 1 month after laser treatment. TEM revealed a significant reduction of BrM thickness in laser-treated eyes of all four groups compared to their fellow control eyes.

Conclusions: TS-R reduces BrM thickness in AMD mouse models ApoE−/− and NRF2−/− without damage to the neuroretina. It may become a prophylactic or even therapeutic treatment option for dry AMD.

Translational Relevance: TS-R may become a prophylactic or even therapeutic treatment option for dry AMD.

Introduction

Age-related macular degeneration (AMD) is a progressive degenerative disease of choroid, Bruch’s membrane (BrM), retinal pigmented epithelium (RPE), and neuroretina. It commences after the age of 50. In the industrialized world, early stage AMD affects one-third of the population at 75 years of age or older—advanced AMD more than 7% of the population at that age.1 AMD is the cause of approximately 50% of severe visual impairment and the most common cause for legal blindness of the elderly in the developed world.2 The need for early and prophylactic AMD therapies to prevent the occurrence of advanced AMD, often followed by legal blindness, is unmet.

AMD can be divided into two different forms, the slowly progressive nonexudative (dry) form and the fast-progressive exudative (wet) form. Dry AMD is characterized by alterations of the RPE leading to
pigment clumping and by metabolic deposits within the RPE-BrM-complex called drusen. Dry AMD may develop to an advanced stage characterized by RPE- and neuroretinal atrophy or it may evolve to wet AMD, which leads to legal blindness quickly in its natural course. Wet AMD is characterized by the appearance of pathological neovascularization of the choroid.

The multifactorial pathology of AMD is not entirely understood. Besides age, genetic variations involving, for example, complement factor H (CFH), complement factor I (CFI), tissue inhibitor of matrix metalloproteases (TIMP3)\(^5\) and factors like high caloric nutrition, lack of physical exercise, oxidative stress, and smoking\(^6\) have been shown to promote AMD. The current concept of the underlying molecular mechanisms of AMD is based on four assumptions: impairment of lipid metabolism, extracellular matrix modifications, inflammatory processes, and altered angiogenesis.

Lipid metabolism of the RPE is essential to provide sufficient nutrients and phospholipids consumed by photoreceptors. The metabolism of lipophilic substances leads to an accumulation of esterified cholesterol within the RPE, which is recomposed with apolipoprotein (Apo) B produced by the RPE itself. ApoB containing high density lipoprotein (HDL) like lipoproteins are excreted basolaterally to BrM and under physiological conditions absorbed by choroid capillary plasma. In AMD, this metabolic pathway is impaired and lipophilic metabolites accumulate in the RPE-BrM-complex.\(^7\)

The multilayered BrM is composed of extracellular matrix and represents the compound basement membranes of RPE and choriocapillaris. It moderates the exchange of nutrients and gas between the choroid and RPE. The homeostasis of this collagenous mesh is mediated by matrix metalloproteases (MMP) and their antagonists tissue inhibitors of metalloproteases (TIMP) secreted mainly by the RPE and to a lesser extent by endothelial cells of the choroid.\(^8\)–\(^11\) In AMD, the homeostatic processes of degradation and rebuilding of BrM are malfunctioning. This leads to an increase in BrM thickness enhanced by metabolic deposits.

Deposits within the RPE-BrM-complex and extracellular matrix modifications under the influence of oxidative stress or genetic predisposition (e.g., malfunction of the complement regulator CFH) may trigger inflammatory processes.\(^12\)–\(^14\) Homeostatic para-inflammatory mechanisms are abandoned in AMD and features of inflammation become evident. Components of the complement system, inflammatory cytokines (interleukin [IL]-6, tumor necrosis factor [TNF]-\(\alpha\)), macrophages, neutrophils, and other immune cells can be found within and around BrM\(^14\)–\(^17\) and especially in drusen.\(^18\) Parts of the innate immune system as well as external factors lead to an increase in oxidative stress and thereby to oxidation of the abundant lipids in and around BrM forming, for example, \(\alpha\)-(2-carboxyethyl)pyrrole (CEP).\(^19\)

Under physiological conditions angiogenesis underlies a homeostasis of pro- and antiangiogenic factors. In AMD, the RPE is stressed by malnutrition and hypoxia deriving from a thickened BrM that inhibits nutrient and gas exchange. Additionally, the RPE is stressed by oxidative processes and stimulated by proinflammatory cytokines. All of this is leading to an increased expression of vascular endothelial growth factor (VEGF), evoking vessel growth originating from the choroid.\(^20\)–\(^22\)

In summary, one of the main features of AMD is a pathologically thickened BrM. A therapeutic means to reduce BrM thickness could therefore be a promising option to cure early stages of AMD and prevent disease progression. To date, there is no effective treatment that addresses early AMD-related mechanisms with a preventive approach.

Laser irradiation of the retina has various effects on RPE and neuroretina. Former studies have shown that laser photoagulation of patients with early AMD aimed on drusen may lead to degradation of these metabolic deposits. However, this method has no influence on visual outcome and does not alter disease progression.\(^23\) One of the main reasons for the therapeutic failure of supra-threshold laser photoagulation for the treatment of AMD is retinal necrosis.\(^23\)–\(^25\) The anatomical region of pathological processes in patients with AMD is the macula and there especially the fovea region. Supra-threshold laser photoagulation is prohibited in the fovea region, since thermal necrosis of the fovea would lead to irreversible vision loss or even legal blindness. Since supra-threshold laser photoagulation is no option for the treatment of early AMD, sub-threshold laser therapy has been of recent interest.

The photomechanical selective retina therapy (SRT) and retinal rejuvenation therapy (2RT\(^\text{e}\)), as well as the photothermal transpupillary thermotherapy (TTT) and thermal stimulation laser therapies (like thermal stimulation of the retina [TS-R] and nondamaging retinal laser therapy [NRT]), are the known laser treatment modalities that neither affect neuroretinal anatomy nor function.
SRT uses a train of microsecond laser pulses that leads to the formation of microbubbles at the intracellular RPE melanosomes, causing a selective disruption of RPE cell membranes. RPE cell death is followed by migration and proliferation of the remaining RPE cells until the RPE monolayer is restored. SRT has shown therapeutic effect on central serous chorioretinopathy and diabetic macular edema, as well as other entities that lead to the formation of subretinal fluid or retinal edema in clinical pilot trials. SRT is not investigated here.

2RT is a nanopulse laser technology that leads to RPE cell death, unevenly distributed within the laser spot, without damage to the neuroretina. It has been shown that 2RT may alter AMD-like alterations in the ApoE−/− model, as well as in humans to some degree. It is currently under investigation within the scope of a phase 3 clinical trial. This laser modality is also not investigated here.

TTT is a laser modality that increases RPE temperature by the application of millimeter-wide spots with long exposures (60 seconds) to the macula. It is hypothesized that it increases the temperature in the irradiated area, especially affecting dividing cells, as seen in wet AMD choroidal neovascularisation (CNV) formation, leading to a decrease of CNV as therapeutic approach. However, repeatedly observed damage to neuroretinal cells, due to lack of power control mechanisms, TTT has not been under investigation anymore.

Thermal stimulation therapies (TS-R and NRT) are continuous wave (cw) or micropulse laser thermal stimulations that lead to a sublethal temperature increase within the RPE without induction of necrosis or apoptosis in RPE or neuroretina. They have shown therapeutic effect in diabetic macular edema and central serous chorioretinopathy. Our preliminary previous data have shown that TS-R induced hyperthermia influences the RPE secretion of AMD-relevant enzymes and signal proteins such as MMP, VEGF, and pigment epithelium-derived factor (PEDF) in porcine organ cultures. Active MMP-2 is upregulated, VEGF downregulated, and PEDF upregulated (Richert E, et al. under review). In vitro studies demonstrated that treatment of human BrM with activated MMPs leads to an enhancement of transport capacity. We hypothesize that TS-R may lead to a restoration of the homeostasis of MMPs and thereby to a restoration of BrM, possibly facilitating trans-BrM metabolic exchange and RPE/photoreceptor nutrition. It might therefore be a therapeutic option for dry AMD.

To test this hypothesis, we evaluated the effect of TS-R on BrM thickness in two AMD mouse models where untreated fellow eyes served as controls.

Two knockout mouse models were chosen to cover the four assumptions that underlie the molecular mechanisms of AMD: impairment of lipid metabolism, remodeling of extracellular matrix, inflammatory processes, and altered angiogenesis.

Apolipoprotein E (ApoE) knockout mice have a reduced catabolism of triglycerides and low-density lipoproteins and are therefore hyperlipidemic and hyperlipoproteinemic. Lipids are known to accumulate in blood vessels and in BrM in these animals leading to an increase in BrM thickness and remodeling of its structure. This strain is known to show AMD-like alterations of the Choroid/BrM/RPE Complex from 2 months of age.

Nuclear factor E2-related factor 2 (Nrf2) knockout mice, on the other hand, lack one of the most important transcription factors of antioxidative processes. Antioxidant response elements (AREs) and drug metabolizing enzymes (DMEs) are downregulated in these mice leading to an increase in cellular stress and accumulation of oxidized cell metabolism products within the RPE and in BrM. This leads to secondary RPE degeneration, inflammatory processes, BrM thickening, and in elderly mice CNV formation. AMD-like alterations have been described from 9 months of age.

Methods

AMD Mouse Models

Both knockout strains (ApoE−/− and NRF2−/−) were purchased from the Jackson Laboratories (Bar Harbor, ME) and housed and bred at the local university animal care unit. The homozygous genotype was confirmed by polymerase chain reaction (PCR) from tail clips. The wild type C57BL6/J control mice were obtained from the local animal care facility. Mice were kept on a regular 12-hour night and day cycle and fed standard murine diet and water ad libitum. All animal experiments were conducted in accordance to the EU directive 2010/63/EU for animal experiments. They were approved by the animal ethics and welfare committee (approval number: V 242-7224.121-12 [61-5/14]) located at the ministry of energy transition, agriculture, environment, and rural areas in Schleswig-Holstein according to German federal and European law. Animal experiments adhere to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.
Animal Maintenance and Anesthesia During Experiments

All examinations and laser treatment were conducted under general anesthesia. Deep anesthesia was induced by an intraperitoneal injection of a mixture of 0.05 mg/kg body weight fentanyl (Braun, Melsungen, Germany), 5.0 mg/kg body weight midazolam (Hameln Pharma, Hameln, Germany), and 0.5 mg/kg body weight medetomidine (CP-Pharma, Bergdorf, Germany).

The animal was then placed on a rigid examination platform, and body temperature was maintained within normal limits using a heating mat. Pupils were dilated by 0.025 mg/mL phenylephrine and 0.05 g/mL tropicamide (UKSH Pharmacy, Kiel, Germany), and eyes were covered with a protective moisturizing gel (2% Methylcellulose Methocel®, Puchheim, Germany). After each examination, the anesthesia was antagonized by a mixture of 1.2 mg/kg body weight naloxone (Braun), 0.5 mg/kg body weight flumazenil (Inresa GmbH, Freiburg, Germany), and 2.5 mg/kg body weight atipamezole (Orion Pharma, Espoo, Finland). Anesthesia was uneventful in all mice. Animal wellbeing was evaluated by a standard score sheet and was uneventful in all included mice. After the final examination, animals were euthanized by cervical dislocation at the day of enucleation under deep anesthesia.

Examinations

All examinations were conducted under general anesthesia. All mice were examined by funduscopy using a contact fundus camera (MICRON III, Phoenix Research Labs, Pleasanton, CA). The integrity of the retina was assessed, the number of drusen-like retinal spots (DRS) was counted, RPE atrophy evaluated, and CNV noted.

Optical coherence tomography (OCT) (small animal OCT, Thorlabs, Lübeck, Germany) was applied via contact optics to evaluate the retinal structure, confirm retinal integrity after laser treatment, and to confirm CNV. After laser treatment, it was used to investigate anatomical integrity of retinal layers.

Fluorescein (10% Fluorescein, 60 mg/kg body weight in 100 µL saline intraperitoneally; Alcon Pharma, Freiburg, Germany) angiography (FLA) (MICRON III) was performed after laser treatment to evaluate vessel integrity, to detect vessel leakage, and to confirm possible CNV formation.

All examinations were repeated at the day of enucleation, thus 1 month after laser treatment.

Figure 1. Laser spot titration and distribution. (A) Laser spots were titrated in the peripheral murine retina (red box) starting from above threshold (from left to right: first white burn) to barely visible lesions (third barely visible burn). The red spot is the aiming beam. TS-R laser treatment was carried out further centrally (red line). All treatment spots were funduscopically invisible. In OCT images, titration lesions were visible (B), and TS-R treated areas showed no anatomical alterations in OCT (C). TS-R treatment spots were distributed uniformly across an optic nerve centered approximately 50° field of view at two-spot interspot spacing, sparing optic nerve disc and retinal vessels (D).

Laser Treatment

For TS-R a frequency doubled Neodym-Vanadate-Laser (Nd:VO₄) experimental laser (Carl Zeiss Meditec AG, Jena, Germany) with a wavelength of 532 nm was used. The light was coupled to an optical multimode fiber of square cross section with a 70 × 70 µm² core profile. The laser light was applied via contact laser-injector (Phoenix Research Labs) attached to the MICRON III camera. The pilot laser was controlled visually via MICRON III live fundus imaging.

Spot size was determined by the size of the laser fiber and laser injector, fixed to 50 µm and could not be changed in size. Duration of irradiation was fixed to 10 ms. The intended stimulating effect was titrated visually by decreasing power at the peripheral retina from a clearly visible white burn at higher power to a barely visible spot at lower power (see Fig. 1). The barely visible burn was classified threshold. Power
was reduced by 70% to determine stimulating laser power similar to the work of Lavinsky and Palanker.40 Laser spots were distributed uniformly across the retina at two spots interspot spacing to an optic disc centered approximately 50° field of view. No laser spot was aimed on vasculature.

**Transmission Electron Microscopy (TEM)**

Enucleated murine eyes were fixated in 4% glutaraldehyde (Merck, Darmstadt, Germany) for 24 hours. Afterwards, the anterior segment was removed under stereomicroscopic (Stemi SV6, Carl Zeiss Meditec AG) control. The posterior eye cup was then embedded into araldite as follows: it was watered in phosphate-buffered saline (PBS; Serva, Heidelberg, Germany) overnight and put into 2% osmium tetroxide (Merck) for 2 hours the next day. Afterwards, eyes were dehydrated in increasing ethanol series at room temperature (70% for 30 minutes, 80% for 40 minutes, 90% and 99% for 50 minutes) and then incubated in propyleneoxide (Merck) 2 × 10 minutes each. A mixture of 91 mL araldite-M (Sigma, Steinheim, Germany) and 84 mL dodecenylsuccinic anhydride (epoxy embedding medium hardener; Sigma) was augmented by 3% araldite M accelerator (Sigma). This mixture was stirred 1:1 with propylene oxide, and the eyes were placed into this medium at room temperature overnight. The next day, araldite combined with 2% araldite M accelerator was added for 4 hours. Eyes were then incubated for 2 hours at room temperature and then hardened at 48°C overnight. Next day temperature was increased to 60°C for another 48 hours.45

Araldite embedded eyes were cut by microtome to sections of 100 nm following a standardized protocol. The first set of three sections was conducted inferior to the rim of the optic nerve, the second set crossed the optic nerve, and the third set of sections was conducted superior to the rim of the optic nerve (see Fig. 2). These sections were contrasted by lead citrate (Merck) and uranyl acetate (Merck) and then examined by TEM Zeiss EM 900 (Carl Zeiss Meditec AG). To eliminate observer bias, the calculation of BrM thickness was done in a standardized and blinded manner by a single trained biologist. From each set of three sections, the first section was examined (two sections in case of damage) taking three pictures at defined spots in proximity to grid bars anterior to the optic nerve and three pictures posterior to the optic nerve. The first picture was always taken strictly behind the first grid bar adjacent to the start of regular RPE morphology in proximity to the optic nerve cavity (where there is no RPE). The second picture was taken strictly before the second grid bar, the third picture strictly behind the second grid bar. Three measurements per picture were carried out. Areas of intercapillary pillars were left out. In total, 54 measurements per eye determined the thickness of BrM (Fig. 2).

**Genotyping**

The validity and homogeneity of the intended genotype as well as screening for retinal degeneration was reduced by 70% mutation of Crumbs-homologue 1 (CRB1<sup>rd8</sup>), known to confound retinal phenotype, was determined by PCR from tail clips. According to Mehalow et al.46 for CRB1<sup>rd8</sup> the following set of primers was used: mCRB1 mF2 (5'-GCCCTGTTTGCATGAGGAAACTTTGGAAGACAGCTA-CAGTTCTTCTG-3'), mCRB1 mF1 (5'-GTGAAGACAGCTA CAGTTCTGATC-3'), and mCRB1 mR (5'-GCCCATTTGCACACTGTAGAC-3'). Nrf2 mice had been genotyped since breeding was more productive in homozygous Nrf2<sup>−/−</sup> + heterozygous Nrf2<sup>−/+</sup> mice, but only homozygous Nrf2<sup>−/−</sup> could be included into the study. According to manufacturer’s specifications, for Nrf2 the following primers were used: Nrf2 c_forward (5'-GCCTGAGAGCTGTAGGGCCC-3'), Nrf2 Wt_reverse (5'-GGAAATGGAAAATAGCTCCTGCC-3'), and Nrf2 Mut_reverse (5'-GACAGTATCGGCCTCAGGAAA-3'). ApoE mice were not genotyped before experiments, because all parental mice were homozygous as specified by the vendor.

After PCR, an agarose gel electrophoresis was carried out. Since two makeup were prepared, for CRB1<sup>rd8</sup> two possible genotypes were expected: wild-type (220 base pairs) and mutant (244 base pairs). For Nrf2, one makeup was prepared, hence three possible genotypes were expected: wild-type (262 base pairs), heterozygous (262 and 400 base pairs), and homozygous (400 base pairs).

**Experimental Protocol**

First, the validity of the two knockout strains as representative of AMD was to be confirmed. Untreated C57BL/6J mice of 5 and 13 months of age were examined clinically (funduscopy, OCT, FLA) and by TEM. ApoE<sup>−/−</sup> of 5 and Nrf2<sup>−/−</sup> of 13 months of age were also examined as untreated control. For the untreated controls both eyes were included. Then, experimental treatment groups had to be defined. From the literature, it is known that...
ApoE−/− mice from the age of 2 months and most ApoE−/− mice at the age of 8 months, as well as NRF2−/− mice from the age of 9 months and most NRF2−/− mice at the age of 12 months show AMD-like alterations of the choroid/BrM/RPE complex.43,44 For both strains, an early (4 months old ApoE−/−, 9 months old NRF2−/−) and a later stage of AMD development (8 months old ApoE−/−, 12 months old NRF2−/−) was chosen. Since the treated mice were enucleated 1 month after TS-R, at the age of 5 and 9 months for the ApoE−/−, and 10 and 13 months for the NRF2−/−, untreated controls were also chosen at comparable ages.

One randomized eye of each mouse in the four different treatment groups (two of different age, two of different strain) of five mice each was treated by TS-R; the fellow eye served as control (see Table 1).

Figure 2. TEM. The anterior segment of the enucleated and fixated eye ball (a) was embedded in araldite and sectioned at three optic nerve disc (ON) centered levels (b). The sections were placed on a copper grid with an intergrid-distance of 150 μm. Grid bars were counted starting from the first grid bar that covered a piece of retina that showed all outer retinal layers in the right anatomical order (the second grid bar shown in scheme c). Therefore, the maximum distance of picture one from optic disc was 300 μm, the minimum distance 150 μm. All pictures were taken within an optic disc centered radius of 450 μm. Three pictures (12,000-fold magnification) to both sides of the optic nerve of each section were taken (d) always in front of the first and in front of and behind the second grid bar (black middle bar) generating a total of six pictures per section. Within each picture, three measurements of BrM thickness were conducted (e, f). The thereby determined BrM thickness is a result of 54 measurements per eye. An overview picture (3000-fold magnification) of each level was taken to examine ultrastructural integrity of RPE before BrM thickness was measured in higher magnification.
Table 1. Number of Mice, Eyes, and Measurements Included in Each Group

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<th>Strain</th>
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<th>Control (c)</th>
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<tr>
<td>ApoE−/−</td>
<td>5</td>
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<td>NRF2−/−</td>
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In the NRF2−/− 13-month group, one mouse died preterm. Above, untreated controls, both eyes were included; Below, mice from laser treatment groups, untreated fellow eyes are controls.

Statistics and Power Calculation

The primary endpoint of the study was BrM thickness. Secondary endpoints were alterations of the AMD-like phenotype and ultrastructural changes of the RPE. The comparison of the laser effect was done intraindividually. Therefore, numbers could be kept low. Since the presented data were a pilot trial with unknown effect, no power calculation was done.

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Statistics and Power Calculation

The primary endpoint addressed intraindividual comparison of BrM thickness in treated versus untreated fellow eyes. TS-R power titration in the retinal periphery produced barely visible spots at 10.5 mW ± 2.0. Power was then reduced by 70% to a mean of 3.8 mW ± 1.4. A mean of 91.5 ± 9 spots was placed uniformly across the optic disc centered approximately 50° field of view fundus (see Fig. 2). TEM revealed a significant reduction of BrM
thickness in 18/19 laser treated eyes (one mouse died preterm) compared to their fellow control eyes ($P < 0.0001$; Fisher) 1 month after laser treatment (see Fig. 6). Five-month-old ApoE−/− control eyes had a mean BrM thickness of 586 nm +/- 12.9 SEM, the TS-R laser treated fellow eye 525 nm +/- 11.4 SEM ($P = 0.0005$; paired $t$-test). Nine-month-old ApoE−/− control eyes’ mean BrM thickness was 603 nm +/- 12 SEM, mean BrM thickness of the TS-R laser treated fellow eyes was 527 nm +/- 12 SEM ($P < 0.0001$). Mean BrM thickness of 10-month-old NRF2−/− control mice was 666 nm +/- 13.3 SEM, the TS-R laser treated fellow eyes had a mean BrM thickness of 598 nm +/- 13.1 SEM ($P = 0.0003$). Thirteen-month-old NRF2−/− control eye mean BrM thickness was 662 nm +/- 12.2 SEM, laser treated fellow eyes had a mean BrM thickness of 593 nm +/- 11.8 SEM ($P < 0.0001$).

Influence of TS-R on Disease Phenotype

Subthreshold TS-R laser spots were not visible by fundus imaging, OCT, and FLA, 2 hours or 1 month after laser treatment. Only threshold titration lesions were visible.

Since CRB1$^{rd8}$ might have influenced the clinical phenotype of NRF2−/−, DRS were counted, but not compared statistically for this strain. Only DRS found in ApoE−/− were compared statistically. There was no significant change in the number of DRS of the treated eye before TS-R compared to the treated eye after TS-R and compared to the control eyes (see Table 2). However, the fellow control eyes of 4-month-old ApoE−/− mice had a significantly increased ($P = 0.03$) number of DRS when compared to 5 months of age.
Figure 4. TEM (12,000-fold magnification) of RPE/BrM/choroid complex pathologies in AMD mouse models. Numbers indicate BrM layers: 1, basal lamina of RPE; 2, inner collagenous layer; 3, elastic layer; 4, outer collagenous layer; 5, basal lamina of choriocapillaris. a single asterisk shows basal lamina splicing, double asterisk shows endothelial thickening, both typically found in AMD, b asterisk indicates outer collagenous layer deposits, and c asterisk indicates the formation of vacuoles within the outer collagenous layer.
Influence of TS-R on RPE Integrity and Ultrastructural Changes

RPE cells are defined as post mitotic polarized epithelial cells with apical melanosomes and long microvilli interdigitating with photoreceptor outer segments. They have little basal pigmentation and short basal microvilli connecting with BrM (Fig. 7B). There was no significant change between treated and untreated eyes in terms of general ultrastructural integrity in 5- and 9-month-old ApoE−/− and 10-month-old NRF2−/−. However, RPE cells with multiple vacuoles and reduced length of apical microvilli were more common (P = 0.04) in the fellow untreated eyes (10 of 12 levels, Fig. 7C) compared to TS-R treated eyes (4 of 12 levels, Fig. 7D) in 13-month-old NRF2−/− mice.

Discussion

We present here that TS-R reduces BrM thickness, known to be pathologically enlarged in AMD. A thickened BrM leads to impaired gas and nutrient exchange, likely being one of the main causes for AMD. We hypothesize that a reduction of BrM thickness may facilitate gas and nutrient exchange and thereby prevent AMD disease progression or even cure the disease. Since neuroretina is left intact, TS-R can even be applied to critical regions of the retina, like macula and could therefore be used as an early treatment for AMD.

Both the ApoE−/− and NRF2−/− mouse models are adequate for the study of AMD therapies, since both models show a thickened BrM with AMD-like ultrastructural BrM alterations. Also, the clinical phenotype mimics AMD with DRS and pigment alterations. With ApoE−/− mice reflecting the hyper-
lipidemic/hyperlipoproteinemic western lifestyle and with NRF2 reflect increased oxidative stress, two of the main driving factors of AMD are present in our models. These two models together include all four main assumptions underlying the pathomechanism of AMD, altered lipid metabolism, changes of the extracellular matrix, inflammation, and altered angiogenesis. Our examinations allow for a statement on the effect of TS-R on lipid metabolism deposits (like drusen) and extracellular matrix. Angiogenesis and inflammation were not in the focus of the study. The interpretation of phenotypical AMD-like retinal alterations may be confound by CRB1rd8. It is known that this mutation mimics AMD-like pathology clinically, for example by an abundance of DRS. CRB1rd8 linked DRS are neuroretinal lesions and not, like in AMD, lesions within the RPE/BrM/choroid complex. This mutation is common in BL/6N background and uncommon in BL/6J background. Since both strains used in this experimental setting are created on BL/6J background, we did not expect to find CRB1rd8. However, our analysis showed that all NRF2 had a homozygous CRB1rd8 mutation, none of the other strains were.

<table>
<thead>
<tr>
<th></th>
<th>DRS Pretreated, n</th>
<th>DRS Posttreated, n</th>
<th>DRS Prefellow, n</th>
<th>DRS Postfellow, n</th>
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<tr>
<td>ApoE−/− 4 mo</td>
<td>4 +/- 4</td>
<td>7 +/- 3</td>
<td>1 +/- 2</td>
<td>5 +/- 2</td>
</tr>
<tr>
<td>ApoE−/− 8 mo</td>
<td>3 +/- 5</td>
<td>5 +/- 5</td>
<td>5 +/- 5</td>
<td>8 +/- 8</td>
</tr>
<tr>
<td>NRF2−/− 9 mo</td>
<td>29 +/- 25</td>
<td>34 +/- 31</td>
<td>25 +/- 60</td>
<td>34 +/- 56</td>
</tr>
<tr>
<td>NRF2−/− 12 mo</td>
<td>31 +/- 28</td>
<td>26 +/- 29</td>
<td>22 +/- 18</td>
<td>25 +/- 19</td>
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</tbody>
</table>

There were no significant differences between the number of DRS before laser treatment compared to 1 month after treatment. However, in the ApoE−/− 4-month-old fellow control eyes (bold letters), the number of DRS was significantly (*P* = 0.03) increased 1 month after treatment (5 months of age) compared to the number of DRS before treatment.

Figure 7. Ultrastructural anatomy of RPE cells. (A) 1100-fold magnification of a TS-R lasered 5-month-old ApoE−/− eye 1 month after treatment showing physiological anatomy. Photoreceptor outer segments interdigitate with long apical microvilli (MV) of RPE cells. Melanosomes (red asterisk) are more common in apical MV and apical cell body. Basal microvilli are free of melanosomes and connect with BrM. Choriocapillaris (CC) is located posterior to BrM. (B) 3000-fold magnification of a TS-R lasered 5-month-old ApoE−/− eye 1 month after treatment showing physiological anatomy. (C) 3000-fold magnification of the untreated fellow control eye of a TS-R lasered 1-month-old NRF2−/− eye 1 month after treatment. Multiple vacuoles (red asterisk) within the RPE cell and a reduced number and length of apical microvilli can be seen. (D) 3000-fold magnification of the TS-R treated eye of C showing physiological anatomy.
affected. The experiments conducted here are translational with focus on BrM thickness. The AMD-like pathology of the RPE/BrM/choroid complex should not be affected by CRB1rd8. Changes due to the mutation are expected within neuroretinal layers. It may though affect the clinical phenotype. Indeed, the variability and in some individuals the abundance of DRS in NRF2−/− suggests CRB1rd8 involvement (Fig. 3). To exclude any confounder, DRS count in NRF2−/− mice was excluded from statistical analysis. CRB1rd8 is unlikely to be a confounder concerning the primary endpoint of the study.

We can clearly demonstrate that TS-R leads to BrM thickness reduction in both AMD mouse models, independently of age, providing evidence for the efficacy of the treatment, even after one laser session. The pilot study presented here cannot make any statement concerning dose–response relations. This analysis needs to be done in future studies.

Besides the above described influence on BrM thickness as primary endpoint, we also examined the effect of TS-R on the phenotype of AMD-like alterations in our mouse models as secondary endpoint. We did not see any influence of the single alterations in our mouse models, independently of age, providing evidence for the secondary effect of TS-R on the phenotype of AMD-like thickness as primary endpoint, we also examined the thickness of BrM and BrM/choroid organ cultures TS-R induces MMP secretion as latent enzymes with intracellular or membrane-associated activation preforms to fully active enzymes. Active enzymes can degrade all components of extracellular matrix, which stimulates cell migration and matrix turnover. This could lead to a reduction of BrM thickness. Thermal stimulation induces the protective chaperones heat shock proteins (HSP) in rabbit models. HSPs not only prevent cells from apoptosis by protein refolding mechanisms, but also induce anti-inflammatory processes. A moderate temperature increase to a maximum temperature of 43°C induces antioxidant cell-protection mechanisms increasing the reduced glutathione (GSH)/oxidized glutathione (GSSG) ratio in RPE cell cultures. TS-R reduces VEGF expression and induces PEDF expression leading to an antiangiogenic milieu (Richert E, et al. IOVS. 2016;57:ARVO E-Abstract 4442). It is unknown whether thermal stimulation affects lipid metabolism. The above-mentioned mechanisms have been investigated in healthy organisms. It is unknown to what extend AMD pathomechanisms are addressed by TS-R in disease. The molecular mechanisms of TS-R should be assessed further.

TS-R has not shown clinical efficacy in AMD yet. It is for the above-mentioned molecular changes induced by TS-R and based on the data presented here—a promising preventive or therapeutic approach for AMD treatment. A major concern though is the reliability of the nondamaging character. Especially if TS-R is considered a preventive measure in the critical macular region, the safety of treatment is of utmost importance. A temperature of approximately 45°C
should likely be the goal to create the wanted biological effect, without cell damage. Spot size in this study was fixed to 50 μm, due to fiber size and laser-injector device. However, since future treatment is aimed for the macula, a small spot size is reasonable to enable a well-controlled application of laser spots in proximity to the fovea region. Also, divergent power within a spot due to the speckle pattern is reduced to a minimum within small spots enabling a homogenous distribution of power within a spot. Irradiation time was fixed to 10 ms. The Arrhenius integral suggests that overall tissue damage is an integration of temperature over irradiation time. Hence, to reliably reach a desired degree of damage (for TS-R a temperature increase without damage is wanted) one must reach a certain temperature integral for a given irradiation time. The impact of inter- and intraindividual variances in pigmentation, optical media, and alignment on the extend of laser-induced temperature increase within RPE cells and adjacent tissue is the bigger, the longer a certain temperature curve has to be maintained. Therefore, cw laser irradiation time was reduced to a minimum of 10 ms. Following this, laser effect titration was only determined by power, controlled clinically by fundus imaging to detect retinal whitening and controlled by OCT to detect structural changes within the retina. This individual titration accounts for the general nature of optical media and species. An individual automatic dosimetry for mice that accounts for intraindividual changes in fundus pigmentation, optical media, and alignment for every spot is not available and would be difficult to obtain due to the size of a murine eye. Functional testing, like electroretinogram, has not been obtained. This would be a valuable examination for future studies not only to evaluate safety.

Barely visible retinal burns are expected to develop just over 60°C.62–65 This taken, the treatment temperature we achieved after power reduction by 70% was approximately 45.2°C, with uncertainty concerning intraindividual variations. Retinal whitening or structural alterations in OCT imaging were only seen in titration areas and never detected in the treatment areas, thus assuring the nondestructive character. For patient treatment, the best evaluated laser power titration system is Endpoint Management.40 Power is reduced to 30% of the last instantly visible laser spot lesion by an Arrhenius curve based algorithm. This titration modality, however, does not recognize intraindividual topic changes in light absorption capacity leading to possible over- or undertreatment. Other dosimetry systems, like optoacoustic feedback mechanisms titrating each laser spot during the laser treatment, have been investigated experimentally in rabbits and humans with good results.66,67 The only laser therapy that does not affect neuroretina, at least anatomically, known to influence AMD-like alterations in humans is 2RT™. A phase 3 clinical trial is underway. The effects on the choroid/BrM/RPE/ neuroretina complex are similar to the effects we see in TS-R, except for the intended RPE cell death.

The limitations of this study are the small number of animals and the lack of knowledge concerning the number of treatments needed, the timing of the treatment, and technical treatment parameters. The laser treatment was carried out once and BrM thickness was determined 1 month after treatment. From organ culture laser experiments (Richert E, et al. IOVS. 2016;57:ARVO E-Abstract 4442) we know that cell mediator levels are altered during the first days after treatment. It is unknown whether the induction of BrM altering MMP lasts even longer. We hypothesized that a possible effect on BrM might need time to develop. We therefore chose the examination time point of 1 month after TS-R. The presented data cannot give any understanding of the long-term course of disease after treatment, since we only examined at one time-point after treatment. Mouse models have general limitations. Murine eyes do not have a macula, while the anatomical structure of the retina is very similar to human retinal anatomy. And finally, there are no human clinical data on TS-R for the treatment of AMD, although TS-R has shown an effect on other retinal disease, for example, central serous retinopathy, and it proved to be safe in humans.

Despite these limitations, TS-R is a promising therapeutic approach and is worthy of further research.

**Conclusion**

We conclude that TS-R leads to a reduction of pathologically increased BrM thickness in AMD, possibly by the induction of active MMPs. TS-R has the potential to be a novel and safe prophylactic or even therapeutic option for early AMD.

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*JT and ER contributed equally to this work.

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