Arginase 1 promotes retinal neurovascular protection from ischemia through suppression of macrophage inflammatory responses

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Abstract
The lack of effective therapies to limit neurovascular injury in ischemic retinopathy is a major clinical problem. This study aimed to examine the role of ureohydrolase enzyme, arginase 1 (A1), in retinal ischemia-reperfusion (IR) injury. A1 competes with nitric oxide synthase (NOS) for their common substrate L-arginine. A1-mediated L-arginine depletion reduces nitric oxide (NO) formation by NOS leading to vascular dysfunction when endothelial NOS is involved but prevents inflammatory injury when inducible NOS is involved. Studies were performed using wild-type (WT) mice, global A1−/− knockout (KO), endothelial-specific A1 KO, and myeloid-specific A1 KO mice subjected to retinal IR injury. Global as well as myeloid-specific A1 KO mice showed worsened IR-induced neuronal loss and retinal thinning. Deletion of A1 in endothelial cells had no effect, while treatment with PEGylated (PEG) A1 improved neuronal survival in WT mice. In addition, A1−/− KO mice showed worsened vascular injury manifested by increased acellular capillaries. Western blotting analysis of retinal tissue showed increased inflammatory and necroptotic markers with A1 deletion. In vitro experiments showed that macrophages lacking A1 exhibit increased inflammatory response upon LPS stimulation. PEG-A1 treatment dampened this inflammatory response and decreased the LPS-induced metabolic reprogramming. Moreover, intravitreal injection of A1 KO macrophages or systemic macrophage depletion with clodronate liposomes increased neuronal loss after IR injury. These results demonstrate that A1 reduces IR injury-induced retinal neurovascular degeneration via dampening macrophage inflammatory responses. Increasing A1 offers a novel strategy for limiting neurovascular injury and promoting macrophage-mediated repair.

Introduction
Ischemia-induced retinal neurovascular injury is a primary contributor in blinding diseases that affect neonates (retinopathy of prematurity), working age adults (diabetic retinopathy), and the elderly (branch vein occlusion). The retinal ischemia-reperfusion (IR) injury model has been widely used to study the mechanisms of neurovascular injury in these and other diseases of the central nervous system (CNS) such as stroke.1–4 Therefore, it provides an excellent model to study the neurovascular damage characteristic of many CNS disorders. The lack of understanding of the mechanisms of IR injury-induced neuronal and vascular injury is a critical barrier for developing clinically effective treatments for these conditions.
Arginase has two isoforms, arginase 1 (A1) and arginase 2 (A2)\(^9\). A1, the cytosolic isoform, is strongly expressed in the liver, where it is the central player in the urea cycle\(^7\). The mitochondrial isoform, A2, is expressed in extracellular tissues, especially the kidney\(^8\). Both isoforms are expressed in the retina and brain\(^9\), and have been linked to CNS diseases\(^19\). A1 is expressed in retinal glia\(^10\). After experimental stroke, A1 has been reported to be stronger in myeloid cells with less expression in astrocytes\(^11,12\). A1 and nitric oxide synthase (NOS) enzyme compete for their common substrate the semi-essential amino acid l-arginine\(^13\). A1 upregulation can lead to suppression of nitric oxide (NO) formation by endothelial NOS (eNOS) resulting in superoxide production, endothelial dysfunction, platelet aggregation, and leukocyte activation and attachment to the vessel wall\(^13\). However, A1 expression in “M2-like” anti-inflammatory myeloid cells is thought to reduce NO production by iNOS, and thus can dampen oxidative stress and inflammation\(^15,16\). Interestingly, the number of A1\(^+\), Iba1\(^+\) macrophages/microglia is correlated with post-stroke neuron survival and recovery in mice\(^11\). Recent studies have shown that A1 is expressed exclusively by infiltrating myeloid cells and not by microglia after CNS injury\(^17,18\).

We have previously shown that A2 plays a deleterious role in retinal IR injury\(^19\). Moreover, retinal IR injury is associated with increased expression of A2 and iNOS, and decreased A1\(^19\). While A1 is a marker for M2 macrophages and is known to improve tissue repair, its role in macrophage polarization and neurovascular damage after CNS IR injury has not been studied\(^10\). Here we examined for the first time the role of A1 in retinal IR injury using mice with global and cell-specific A1 deletion. We also tested the therapeutic potential of PEGylated A1 (PEG-A1, a drug form of A1 that is currently under investigation as cancer therapy\(^20–24\)) in retinal IR injury.

Results
A1 deletion worsens IR-induced neurovascular degeneration in vivo

We have previously shown that retinal IR injury is associated with decreased A1 mRNA at 3 h\(^19\). In line with this, we found a sustained decrease in retinal arginase activity starting at 3 h after IR injury and up to 48 h (Fig. S1). To study the role of A1 in retinal IR injury, we used heterozygous (A1\(^+/–\)) global KO mice, since homozygous deletion of A1 is postnatal lethal\(^25\). WT or A1\(^+/–\) KO mice were subjected to 40 min of ischemia on the right eye followed by reperfusion as explained in the methods\(^26\). The left eye served as sham control. The retinal IR injury model is associated with both neuronal and microvascular degeneration that are manifested by neuronal loss and acellular capillary formation\(^19\). To evaluate neurodegeneration after IR injury, we labeled WT and A1\(^+/–\) KO retinas with the neuronal marker, NeuN and imaged the surviving neurons in the retinal ganglion cell layer by confocal microscopy\(^19,26\). IR injury reduced NeuN-positive cells in WT mice at 7 days, which was further worsened in A1\(^+/–\) mice (Fig. 1a, b). We next examined microvascular degeneration preparation by preparing retina vascular digest and counting the number of acellular capillaries\(^19,27\). WT IR injured retinas showed a large number of acellular capillaries (\(\approx 150/\text{mm}^2\) empty basement membrane sleeves, red arrows, Fig. 1c, d) at 14 days after IR injury and this was further increased by \(\approx 50\%\) in A1\(^+/–\) mice.

A1 deletion exacerbates retinal thinning and distortion after IR injury

The IR injury model has been shown to affect the inner retinal layers (ganglion cell layer (GCL), inner plexiform layer (IPL), and inner nuclear layer (INL)) to a greater extent than the outer retina leading to reduced inner retina thickness\(^26,28,29\). In accordance with this, morphometric analysis on hematoxylin and eosin (H&E)-stained WT IR injured retina sections at 7 days showed reduced thickness of the inner retinal layers compared to sham controls. A1\(^+/–\) retinas showed further thinning and distortion compared to WT after IR injury (Fig. 2a, b). This was confirmed by optical coherence tomography (OCT) that showed worsened retinal detachment in A1\(^+/–\) retinas (Fig. 2c).

A1 deletion exacerbates retinal inflammation and necroptosis after IR injury

Next, we examined the underlying mechanism of increased retinal cell death in A1\(^+/–\) mice after IR injury. Various mechanisms of retinal cell death have been described in the retina IR injury model with studies from our lab and others emphasizing a prominent role of programmed cell death by necroptosis (a caspase-independent programmed form of cell death)\(^19,30–35\). Necroptosis is associated with an early increase in cell membrane permeability. We evaluated this through propidium iodide (PI) uptake, which is plasma membrane impermeable and only labels the DNA of dying cells. We observed PI-positive cells in GCL and INL of WT retinas within 6 h following IR injury with more cells in A1\(^+/–\) retinas (Fig. S2).

Unlike apoptosis, necroptosis is associated with release of cellular contents and subsequent inflammatory response. Therefore, we examined the necroptosis marker receptor interacting protein 3 kinase (RIP3) together with other inflammatory markers via western blotting. Western blot analyses showed increases in the stress marker phospho-p38 MAPK in A1\(^+/–\) retinas at 3 h after IR injury as compared to WT retinas. There was also a trend towards an increase in the mitochondrial fission marker, dynamin-related protein (Drp1) but the difference was not statistically significant (Fig. 3a–c). Furthermore, IR injury induced increases in the inflammatory cytokine,
tumor necrosis factor alpha (TNF-α), and RIP3 in A1+/− retinas at 6 h (Fig. 3d–g). The increase in inflammation was associated with increases in oxidative stress. This was shown by increased nitrotyrosine formation (a measure of protein tyrosine nitration via peroxynitrite, which is formed by the reaction of NO with superoxide anion). Albumin extravasation (a measure of vascular permeability) was also increased in A1+/− as compared to WT mice at 48 h after IR injury (Fig. 3h–j).

**Effect of cell-specific A1 deletion on neuronal survival and retinal tissue thinning after IR injury**

It has been shown that retinal IR injury induces macrophage/microglia recruitment and proliferation within 24 h with a peak in cell number at 3–5 days36. In accordance, we have seen an increase in Iba1-positive cells in the retina after IR injury (Fig. S3A). Interestingly, we detected Iba1-positive cells in the vitreous at 48 h after IR injury, suggesting infiltration of systemic monocyte-derived macrophages (Fig. S3B). Building on this and since global A1 deletion showed a worsened retinal IR injury outcome, we next examined the cell-specific role of A1. For this, A1 floxed (loXP) mice were crossed to LysMcre and Cdh5cre transgenic mice to generate mice lacking A1 in myeloid (LysMcre:A1fl/fl) or endothelial (Cdh5cre:A1fl/fl) cells, respectively. Mice with myeloid but not endothelial A1 deletion showed exacerbated neuronal loss at 7 days after IR injury compared to littermate floxed
control (Fig. 4a, b). Moreover, myeloid A1 KO retinas showed more thinning and distortion after IR injury (Fig. 4c, d). Western blotting on retinal lysates from endothelial-specific A1 KO mice showed no difference in albumin extravasation at 48 h after IR injury as compared to floxed controls (Fig. S5). Collectively, these data suggest a major protective role of myeloid A1 and a minimal role of endothelial A1 in retinal IR injury. In line with a reparative role of infiltrating macrophages in the retinal IR injury model, systemic macrophage depletion using clodronate liposomes led to worsened neurodegeneration and retinal hemorrhage after IR injury in WT mice (Fig. S7).

**PEGylated A1 treatment protected retinal neurons and increased microglia/macrophages after IR injury**

To study the effect of increasing A1 levels on neurodegeneration, we used PEG-A1, which is an investigational drug with good safety, pharmacodynamic, and pharmacokinetic profiles in patients. WT mice received intravitreal injection of PEG-A1 (1.7 ng in 1 μl), 3 h before or after IR injury. Either pre- or post-treatment with PEG-A1 improved neuronal survival at 7 days after IR injury. Interestingly, the PEG-A1-mediated neuronal preservation was associated with more retinal macrophages/microglia with elongated morphology as evident by Iba1 staining of retina flat-mounts (Fig. 5a−c).

**A1 deletion augments macrophage inflammatory response in vitro and PEGylated A1 treatment mitigates it**

To further confirm our in vivo data, we tested the role of A1 expression in macrophage inflammatory response in vitro. Peritoneal macrophages isolated from myeloid A1 KO and floxed littermate controls were treated with interleukin-4 (IL-4, 20 ng/ml) or lipopolysaccharide (LPS, 100 ng/ml) for 24 h to generate anti-inflammatory (M-2 like) or proinflammatory (M-1 like) responses, respectively. As expected, IL-4 treatment increased A1 expression in macrophages isolated from control mice.
Upon LPS stimulation, macrophages lacking A1 showed more iNOS expression, TNF-α and inflammasome pathway activation (NLRP3, NFKB, and pro-IL1β) compared to controls. Furthermore, they showed increased NO production in cell supernatant compared to controls as measured by NO analyzer. Collectively, A1 KO macrophages exhibited a more pronounced inflammatory response to LPS stimulation. Cotreatment with PEG-A1 (1 μg/ml) dampened the LPS-induced inflammatory response and reduced NO production in both control and A1 KO macrophages (Fig. 6).

PEGylated A1 rescues LPS-induced mitochondrial dysfunction in macrophages

Next, we examined macrophage metabolic reprogramming, which has been implicated in macrophage polarization and inflammatory response. Seahorse XFe96 analyzer was used to evaluate mitochondrial function by measuring the oxygen consumption rate (OCR). As previously described, LPS stimulation (100 ng/ml) shifted WT bone marrow-derived macrophages (BMDMs) to a more glycolytic phenotype (as measured by increased extracellular acidification rate, ECAR) and decreased mitochondrial respiration parameters (OCR). PEG-A1 (1 μg/ml) significantly inhibited the LPS-induced alterations in mitochondrial function parameters in WT BMDMs (Fig. 7a–g). In addition, staining live BMDMs with the mitochondrial membrane potential sensitive dye, Rhodamine 123, showed mitochondrial fragmentation in response to LPS stimulation which was partially reversed with PEG-A1 cotreatment (Fig. 7i, j). Seahorse analysis of LPS-stimulated A1 KO macrophages showed impaired mitochondrial function as compared to loxP controls (Fig. S8). PEG-A1 treatment of A1 KO macrophages blunted the LPS-induced impaired mitochondrial function (Fig. S9).

NO has been shown to decrease mitochondrial reserve capacity in endothelial cells. We aimed to further examine the link between A1 and mitochondrial respiration in endothelial cells. Bovine retinal
endothelial cells were subjected to oxygen-glucose deprivation (OGD) for 5 h followed by 1 h reoxygenation (R). OGD/R impaired mitochondrial respiration parameters and PEG-A1 treatment (1 μg/ml) improved maximal respiration and spare respiratory capacity (difference between maximal respiration and basal respiration) after OGD/R (Fig. S10). Collectively, PEG-A1 treatment rescued the LPS- and OGD/R-induced...

Fig. 4 Myeloid A1 deletion worsens neuronal loss and retinal thinning after IR injury. a Retinas of mice with myeloid but not endothelial-specific A1 deletion showed worsened neuron loss compared to floxed control at 7 days after IR injury. Scale bar = 100 μm. b Quantification of NeuN-positive cells, n = 9 for A1ffe and 5 for M-A1−/− and E-A1−/− IR, *p < 0.05 vs. A1ffe. c H&E staining at 7 days showed worsened inner retina thinning in M-A1−/− mice compared to control (A1ffe). Scale bar = 50 μm. d Quantification of inner retina thickness, n = 7 for A1ffe IR and 6 for M-A1−/− IR, *p < 0.05. e Optical coherence tomography (OCT) corroborated the H&E results with yellow arrow pointing at retinal detachment.
mitochondrial dysfunction in macrophages and endothelial cells respectively.

**Intravitreal injection of A1 KO BMDMs is associated with worsened neurodegeneration after IR injury**

Finally, we employed intravitreal injections of BMDMs to further examine the role of A1 in the macrophage reparative/damaging functions. Mice were subjected to IR injury and intravitreal macrophage injection was conducted at day 3. Retinas were collected for NeuN staining at day 7. Indeed, A1 KO macrophage injection was associated with worsened neurodegeneration after IR injury compared to loxP control macrophages (Fig. 8).

**Discussion**

In this study, we present evidence on the important role of A1 in macrophage polarization toward a reparative phenotype leading to neurovascular recovery after retinal IR injury. We show that global as well as myeloid-specific A1 deletion worsens retinal IR injury outcomes and macrophages lacking A1 exhibit enhanced inflammatory response. Additionally, PEG-A1 treatment reduces retinal IR injury and dampens the macrophage inflammatory response.

Although retinopathies are diagnosed primarily based on vascular abnormalities, studies have demonstrated that inflammation and neurodegeneration occur before appearance of typical vascular pathology. Our present study in a retinal IR injury mouse model shows that A1 protects against neuronal and vascular injury. This was evident by the worsened neurodegeneration, acellular capillary formation, retinal thinning, and necroptosis in A1+/− mice compared to WT. By contrast, our recent study in A2−/− mice suggests a deleterious role of A2 in retinal IR injury. This could be explained by the differential expression and subcellular localization patterns of the two arginase isoforms. Both of our studies have employed different endpoints to confirm the role of A1 and A2 in retinal IR injury. Other studies have also shown opposing roles of A1 vs. A2 under different disease conditions. In fact, these two isoforms are transcribed from two different genes and appear to function independently in different tissues. However, the possible reciprocal regulation and interaction between A1 and A2 remains to be elucidated.

Necroptosis is initiated in response to a death receptor signaling and upon failure of apoptosis induction. TNF-α binding to its receptor can induce necroptosis with RIP3...
Fig. 6 (See legend on next page.)
being the main executioner. Necroptosis leads to membrane permeability and release of cell components, which causes an inflammatory response. RIP3 can also directly activate the NLRP3 inflammasome pathway. We and others have shown activation of the necroptosis pathway within hours of the IR injury which can induce inflammation and myeloid cell recruitment. Our previous publication 19, and current data show that while A2 deletion reduces IR injury-induced necroptosis in the retina, A1 deletion can augment it.

Our molecular analyses show that A1 deletion is associated with upregulation of iNOS, peroxynitrite, TNF-α, and RIP3. A1 competes with NOS enzymes for their common substrate l-arginine leading to less NO production 13. A1 upregulation can decrease NO production by iNOS in myeloid cells, thus reducing oxidative stress and inflammation 15,16. However, chronic A1 upregulation in endothelial cells can lead to uncoupling of endothelial NOS (eNOS), via reduction of l-arginine, resulting in endothelial dysfunction 14. In our acute model of retinal IR injury, we did not detect any A1-mediated adverse vascular outcomes. In fact, A1+/− mice showed more microvascular degeneration and greater vascular permeability which was not affected by endothelial-specific deletion of A1. Furthermore, PEG-A1 treatment rescued the OGD/R-induced mitochondrial dysfunction in endothelial cells. Collectively, this suggests a vascular protective role of A1 in our acute injury model in young and otherwise healthy mice. Possible adverse vascular effects of A1 treatment in mouse models of endothelial dysfunction will be addressed in future studies.

A1 has been long used as a marker for anti-inflammatory M2-like macrophages; however, its direct functional role has not been examined in CNS diseases. Using myeloid-specific A1 KO mice as well as in vitro macrophage experiments, we show here that this protective role of A1 in IR injury is mediated through myeloid cells. NO induces TNF-α expression in myeloid cells and potentiates its neurotoxicity 45-54. Our results suggest that A1 decreases iNOS-mediated NO production and oxidative stress in myeloid cells leading to decreased TNF-α production and subsequent necroptosis and tissue damage. Moreover, NO has been linked to macrophage metabolic reprogramming to a more glycolytic phenotype and less mitochondrial oxidative phosphorylation through nitrosylation of the electron transport chain complexes 55.

In line with this, our data show that PEG-A1 effectively rescues LPS-induced suppression of mitochondrial metabolism. Interestingly, NFκB activation in response to LPS stimulation was not changed by A1 deletion or PEG-A1 treatment suggesting that A1 regulates iNOS and inflammatory cytokines transcription at a level downstream of NFκB phosphorylation. One possibility is that A1 suppresses inflammatory gene transcription in macrophages through epigenetic modification. A recent report has shown that, putrescine, a downstream product of arginase can suppress M1 inflammatory gene transcription in macrophages through histone modification and altered euchromatin formation 56. Further studies are needed to elucidate the mechanisms by which A1 promote a less inflammatory macrophage phenotype.

Monocyte-derived macrophages infiltrate the CNS after injury and recent reports have examined their role in CNS injury outcome. Two independent studies have shown that brain-infiltrating macrophages after ischemic stroke acquire an M2-like reparative phenotype 57,58. Monocyte-derived macrophages have been shown to prevent hemorrhagic transformation and mediate long-term functional recovery after stroke in mice 59,60. Moreover, brain-infiltrating macrophages reduce lesion volume and neurological deficit in an intracerebral hemorrhage (ICH) mouse model through M2 polarization 61. Interestingly, macrophages have been shown to promote vascular repair after traumatic brain injury in mice, and directly repair cerebrovascular ruptures in zebrafish 62. On the other hand, one report showed that macrophage depletion reduced myelin damage and promoted neurological recovery in a mouse stroke model 65. Our current data show that macrophages play a protective role in retinal IR injury and their depletion further worsens neurodegeneration and hemorrhage. Moreover, we show that A1 is a central player in this protective effect. While our in vitro studies focused on A1-mediated dampening of the macrophage inflammatory response, the in vivo studies...
suggest that A1 can promote a reparative macrophage phenotype as well. The proposed protective role of macrophages in our model could be mediated through clearing of dying cells via phagocytosis and promoting vascular repair. Further studies are warranted to examine these mechanisms.

In conclusion, our study shows that A1 ameliorates the IR injury-induced retinal injury via dampening the macrophage inflammatory response. Enhancing myeloid A1 can be a potential therapeutic approach for treatment of CNS ischemic conditions, especially ischemic retinopathies.
Materials and methods

Mouse retinal IR injury model

All animal procedures were performed in accordance with the Association for Research in Vision and Ophthalmology (ARVO) Statement and adhered to the Public Health Service Policy on Humane Care and Use of Laboratory Animals (revised July 2017). Procedures were approved by the institutional animal care and use committee (Animal Welfare Assurance no. A3307–01). Male mice (10–12 weeks old) were anaesthetized with ketamine/xylazine mixture and the right eye was subjected to IR injury while the left eye served as sham control. IR injury was induced by inserting a needle connected to an elevated saline reservoir into the anterior chamber to raise the intraocular pressure to 110 mmHg for 40 min as previously described19,26. The mice were sacrificed at various times after IR injury to examine different endpoints based on the literature and our previous publications19,26.

Experimental groups

To determine the role of A1 in neurovascular degeneration after IR injury, we used mice with global and conditional A1 deletion: heterozygous A1+/− knockout (KO), endothelial-specific (Cdh5Cre;A1f/f or E-A1−/−) KO, and myeloid-specific (LysMCre;A1f/f or M-A1−/−) A1 KO. A1+/− mice on C57BL6J background were used since A1−/− is postnatal lethal due to hyperammonemia25,65–67. Experiments were conducted on littermate control and KO mice to ensure proper comparison. IR injury was conducted on all mice within a litter during the same day. Mice were selected for surgery in a random order irrespective of the genotype. Cell-specific A1 KO mice were generated by crossing C57BL-6 A1 floxed mice (A1f/f) with Cre-expressing transgenic mice under control of the VE-Cadherin promoter (Cdh5Cre) or lysozyme 2 promoter (LysMCre) to generate endothelial-specific (E-A1−/−) or myeloid-specific (M-A1−/−) mice respectively. We have recently characterized and confirmed the E-A1−/− mice68. Myeloid A1 deletion has been confirmed by tissue immunostaining and western blotting on isolated cells (Fig. S4). Further details are provided in the supplementary data.

Five sets of in vivo experiments were conducted:

Experiment 1: A1+/− KO mice and C57BL6J WT controls were used to examine the effect of whole body deletion of one copy of A1 on IR injury outcome.

Experiment 2: Endothelial-specific (E-A1−/−) and myeloid-specific M-A1−/− mice were used and compared to floxed controls (A1f/f) to examine the cell-specific role of A1.

Experiment 3: WT mice were treated with clodronate liposomes to deplete systemic monocytes/macrophages or control liposomes to examine the role of these cells in retinal IR injury.

Experiment 4: WT mice were treated with intravitreal injection of PEG-A1 (1.7 ng in 1 μl, dose was selected based on preliminary studies) to examine the effect of increasing intraocular arginase on retinal IR injury outcome.

Experiment 5: WT mice received intravitreal injection of A1−/− or A1f/f macrophages to examine the impact of A1 expression in macrophages on IR injury outcome.

Monocyte/macrophage depletion

Two hundred microliters of clodronate or red fluorescent control liposomes (Encapsula Nanosciences) were
injected intraperitoneally 1 day before and at day 3 after IR injury. This achieved ≈80% systemic monocyte/macrophage depletion at day 7 after IR injury (Fig. S6). Clodronate induces apoptosis when the liposomes are engulfed by macrophages while control liposomes were used to control for the induction of macrophage phagocytic activity. Cages were changed every 2-3 days to maintain a clean environment and mice were closely monitored for any signs of infections.

**PEGylated arginase 1 (PEG-A1) treatment**

PEG-A1 (1.7 ng/μl) or phosphate-buffered saline (PBS) was administered under anesthesia via intravitreal injection (in 1 μl volume) using a Hamilton syringe 3 h before or after induction of retinal IR injury.

**Intravitreal macrophage injection**

Cultured BMDMs were suspended in PBS and injected intravitreal (2×10⁵ per 1 μl) on day 3 after IR injury.

**Evaluation of neurodegeneration**

Neuronal degeneration was assessed at 7 days after IR injury as previously described19,26. Eyeballs were collected and fixed overnight in 4% paraformaldehyde at 4 °C, then retinas were dissected into flat-mounts and stained for the neuronal marker, NeuN (Millipore, Cat. # MAB377, Billerica, MA). Four images were taken in the retina midperiphery using a confocal microscope (LSM 510; Carl Zeiss, Thornwood, NY) and NeuN-positive cells were counted using ImageJ software. Results are presented as a percent of NeuN-positive cell numbers in the GCL of the contralateral sham eyes compared to the sham eyes.

**Retinal vasculature isolation and measurement of acellular capillaries**

Vascularity was isolated at 14 days after IR injury via trypsin digestion of retinas that were dissected from overnight fixed eyeballs as previously described19,69. The isolated retinal vasculature was air-dried on silane-coated slides and stained with periodic acid-Schiff and hematoxylin. Acellular capillaries were counted in ten random fields of the mid-retina. The number of acellular capillaries was divided by the field area to get number of acellular capillaries per 1 millimeter square (mm²) of retina.

**Histology and morphometric analysis**

Retinal thickness was assessed at 7 days on anesthetized mice using OCT (the Bioptigen Spectral Domain Ophthalmic Imaging System, SDOIS; Bioptigen Envisu R2200, NC) as previously described70. Retinal thickness was determined by morphometric analysis on H&E-stained retinal frozen cross sections as previously described19,26,70. Inner retina (INL + IPL + GCL) thickness was measured on H&E images at three different distances from optic nerve head using ImageJ software. Averaged retinal thickness was presented as percentage compared to the contralateral sham eyes.

**Western blot analysis**

For in vivo experiments, retinas were collected from the mice, snap-frozen in dry ice, and stored at −80 °C. For analysis of albumin leakage across the blood-retinal barrier, mice were transcardially perfused with PBS to clear blood out of the retina vessels before collection and retinas were processed for western blotting. Retinas were homogenized using a hand homogenizer in RIPA lysis buffer and centrifuged at 20,000 × g to prepare the protein extracts. Protein concentration was measured using Pierce BCA protein assay kit (Thermo Scientific). For in vitro experiments, media were collected and cells were washed with ice-cold PBS then collected in RIPA buffer to prepare the cell lysates. Retinas or cell lysates were run on SDS-PAGE then transferred to nitrocellulose membranes (Millipore, Billerica, MA). The membranes were probed with following primary antibodies prepared in 2% BSA: A1 (Santa Cruz Biotechnology Cat. # Sc-20150, 1:500 dilution), phospho p38 (Cell Signaling Technology Cat. # 4511, 1:500), total p38 (Cell Signaling Technology Cat. # 9212, 1:500), Drp1 (Santa Cruz Biotechnology, Cat. # SC-271583, 1:500), TNF-α (Abcam, Cat. # ab11973), albumin (Bethyl Laboratories, 1:5000), RIP3 (Santa Cruz Biotechnology, Cat. # SC-135170, 1:500), β-actin (Sigma-Aldrich Cat. # A1978, 1:5000), anti-nitrotyrosine antibody (Millipore, Cat. # 05-233, 1:5000), iNOS (Cell Signaling, Cat. # 13120), A2 (Santa Cruz Biotechnology, Cat. # Sc-20151, 1:500 dilution), IL-1β (R&D, Cat. # AF-401-NA), p-NFκB (Cell Signaling, Cat. # 3033, 1:500), T-NFκB (Cell Signaling, Cat. # 4764, 1:500), NLRP3 (Cell Signaling, Cat. # 15101). Secondary antibodies (GE Healthcare) were prepared in 5% milk in 1:2000 dilution. Bands were quantified using ImageJ and normalized to β-actin loading control. For the nitrotyrosine blot, all bands in each lane were quantified except for the thick albumin band (~66 kDa) that appears due to the reaction of the secondary antimouse antibody with mouse albumin in the tissue extracts.

**Isolation and culture of primary macrophages**

**Peritoneal macrophages**

Mice were injected intraperitoneally with 5 ml of 3% Brewer’s thioglycollate medium (Sigma) as described previously27. Mice were sacrificed 3-5 days later and peritoneal macrophages were collected in PBS through peritoneal lavage. Cells were centrifuged and then plated in six-well plates (1 million per well) in DMEM containing penicillin/streptomycin (P/S), and 10% fetal bovine serum (FBS). Medium was changed after 2 h to remove...
nonadherent nonmacrophage cells. Experiments were done on day 2 in DMEM containing 1% PS and 2% FBS.

**Bone marrow-derived macrophages**

Bone marrow cells were isolated and differentiated in vitro into macrophages based on a published protocol. Briefly, both femurs and tibias were harvested and flushed with 20–25 ml sterile PBS using a 27-gauge needle. Flushed cells in PBS were spun down and resuspended in differentiation medium (DMEM high glucose containing 20% FBS, 20% L929 conditioned media, and 1% P/S). Cells were subsequently plated on 100 mm dishes (uncoated, for easier cell detachment). Media was replaced with fresh differentiation media on day 4. On day 7, media was removed and cells were rinsed with sterile PBS twice. Cells were subsequently gently scraped and collected in PBS, spun down, re-suspended in normal growth media (DMEM high glucose containing 20% FBS, and 1% P/S), and then plated in 12-well plates for in vitro polarization or used for in vivo treatment.

**Macrophage polarization**

Cells were stimulated with LPS (100 ng/ml) or interleukin 4 (IL-4, 20 ng/ml) for 24 h to achieve proinflammatory (M1) or anti-inflammatory (M2) phenotype respectively. In some experiments, macrophages were cotreated with LPS and PEG-A1 (1 μg/ml) to examine the effect of PEG-A1 on the LPS-induced macrophage inflammatory response.

**Arginase activity assay**

Arginase activity assay was conducted as previously described. Briefly, the enzyme was activated by heating the lysate or supernatant at 56 °C in 25 mM Tris buffer (pH 7.4) containing 5 mM MnCl₂. L-Arginine hydrolysis was then conducted by incubating 50 μl of the activated samples with 50 μl of 0.5 M L-arginine (pH 9.7) at 37 °C for 60 min. The reaction was stopped by adding 400 μl of acid solution mixture (H₂SO₄:H₃PO₄:H₂O, 1:3:7). The concentration of urea, which is the end product of L-arginine hydrolysis by arginase, was determined after adding 25 μl of 9% α-isonitrosopropiophenone and heating the mixture at 100 °C for 45 min. Urea standards and 200 μl of each sample were transferred to a 96-well plate and read at 540–550 nm in a BioTek microplate reader. Protein concentration in the lysates was determined by a BCA assay (Pierce Biotechnology). Arginase activity was calculated as mmol urea/mg protein and as percent of control.

**Nitric oxide (NO) measurement**

We measured nitrite (NO₂⁻), the stable breakdown product of NO in cell-conditioned medium to reflect NO production by macrophages. The conditioned media were collected at the end of experiments and injected in glacial acetic acid containing sodium iodide. NO₂⁻ is quantitatively reduced to NO under these conditions, which can be quantified by a chemiluminescence detector after reaction with ozone in an NO analyzer (Sievers, Boulder, CO).

**Quantitative RT-PCR**

Total RNA was extracted from macrophages using TRIzol reagent (Invitrogen, CA, USA). RNA was converted to cDNA using M-MLV reverse transcriptase (Applied Biosystems, CA, USA) with SYBR Green dye. Forward primer 5'-GTG GAC GGG TCG ATG TCA C-3', reverse primer 5'-GTT CTC AGC CCA ACA ATA CAA GA-3', and 25 mM glutamine (Gemini, West Sacramento, CA), and 25 mM glucose (Sigma, St. Louis, MO). On the day of the assay, the pH of the media was adjusted to 7.4 ± 0.1, and the Mito stress test was conducted according to the manufacturer's instructions. The concentrations of the injection compounds used were as follows: oligomycin (1 μM), FCCP (1 μM) and rotenone/antimycin A (0.5 μM). The data were normalized to HPRT and the fold change between levels of different transcripts was calculated by the ΔΔCT method.

**Seahorse XFe96 Mito stress test**

Mito Stress test (Agilent, Cat. # 103015-100, Santa Clara, CA) and Seahorse XFe96 (Agilent, Santa Clara, CA) were used to evaluate mitochondrial function as previously described. Briefly, Seahorse 96-well cell culture plates were used for growing the cells. At day 7, BMDMs were seeded at cell density of 40 K/well in the Seahorse cell culture plate in all the wells except A1, A12, H1 and H12 wells which were used as background wells. The plate was left under the hood for 1 h to ensure even distribution of cells, and then cells were checked under microscope and put in the incubator. Cells were maintained to grow in normal complete media (DMEM, 20% FBS, 1% P/S) for 3–5 h. Then, cells were treated with LPS overnight in DMEM media supplemented with 10% FBS. The Seahorse media was prepared according to manufacturer's instructions and supplemented with 4 mM glutamine (Gemini, West Sacramento, CA), 1 mM pyruvate (Gemini, West Sacramento, CA), and 25 mM glucose (Sigma, St. Louis, MO). On the day of the assay, the pH of the media was adjusted to 7.4 ± 0.1, and the Mito stress test was conducted according to the manufacturer's instructions. The concentrations of the injection compounds used were as follows: oligomycin (1 μM), FCCP (1 μM) and rotenone/antimycin A (0.5 μM). The data were collected and analyzed using the Wave software (Agilent).

**Mitochondrial staining**

Vectacell™ Rhodamine 123 (Vector Laboratories) was used to label mitochondria in live cells according to the manufacturer's instructions. Briefly, cells were washed three times with modified PBS containing 1 mM CaCl₂ and 0.5 mM MgCl₂ (PBS⁺) (Thermo Fisher). The cells were incubated with the Rhodamine 123 staining solution...
in 37 °C for 15 min. Then, the cells were rinsed three times with PBS+. Zeiss LSM 780 Inverted Confocal microscopy (Carl Zeiss AG, Oberkochen, Germany) was used to image the live cells using a ×63 lens. Several images were taken randomly to cover the whole field.

**Statistical analysis**

Statistical analysis was conducted using GraphPad Prism 7 software. Values were tested to assess whether they followed a normal distribution by the same software. One-way or two-way ANOVA with post-hoc Tukey multiple comparisons was used to analyze the statistical significance of experimental results in studies of three or more groups. The significance of differences between two groups was determined by Student’s t-test. p < 0.05 was considered significant. Sample size for each experiment was decided based on our previously published work. Outliers were checked by GraphPad online outlier calculator. For in vitro studies, each experiment was performed in triplicates and repeated with at least three different batches of isolated primary cells. Graphs were prepared using GraphPad Prism 7 software, and results were expressed as means ± standard errors of the mean (SEM).

**Disclaimer**

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**Conflict of Interest**

The authors declare that they have no conflict of interest.

**References**

1. Osborne, N. N. et al. Retinal ischemia: mechanisms of damage and potential therapeutic strategies. Prog. Retin. Eye Res. 23, 91–147 (2004).
2. D’Onofrio, P. M. & Koelberle, P. D. What can we learn about stroke from retinal ischemia models? Acta Pharmacol. Sin. 34, 91–103 (2013).
3. Hartsock, M. J. et al. A mouse model of retinal ischemia-reperfusion injury through elevation of intracranial pressure. Jove 113, e54065 (2016).
4. Binet, F. et al. Neuronal ER stress impedes myeloid-cell-induced vascular regeneration through IRE1α degradation of netrin-1. Cell Metab. 17, 353–371 (2013).
5. London, A., Benhar, I. & Schwartz, M. The retina as a window to the brain-from eye research to CNS disorders. Nat. Rev. Neuro. 9, 44–53 (2013).
6. Ash, D. E., Cox, J. D. & Christianson, D. W. Arginase: a binucleic manganese metalozyme. Met. Ions. Biol. Syst. 37, 407–428 (2000).
7. Morris, S. M. Jr. Regulation of enzymes of the urea cycle and arginine metabolism. Annu. Rev. Nutr. 22, 87–105 (2002).
8. Miyanaka, K. et al. Immunohistochemical localization of arginase II and other enzymes of arginine metabolism in rat kidney and liver. Histochem. J. 30, 741–751 (1998).
9. Patel, C. et al. Arginase as a mediator of diabetic retinopathy. Front. Immunol. 4, 173 (2013).
10. Caldwell, R. W., Toque, H. A., Narayanan, S. P. & Caldwell, R. W. Arginase: an old enzyme with new tricks. Trends Pharmacol. Sci. 36, 395–405 (2015).
11. Hamzeti Taj, S., Kho, W., Roii, A., Wiedermann, D. & Hoeib, M. M. ARN-124 induces neuroprotection and functional improvement after focal cerebral ischemia. Biometals 91, 151–165 (2016).
12. Quine, A. et al. Effect of stroke on arginase expression and localization in the rat brain. Eur. J. Neurosci. 37, 1193–1202 (2013).
13. Wu, G. & Morris, S. M. Jr. Arginase metabolism: nitric oxide and beyond. Biochim. J. 336(1 Pt 1), 1–17 (1998).
14. Caldwell, R. W., Rodriguez, P. C., Toque, H. A., Narayanan, S. P. & Caldwell, R. W. Arginase: a multifaceted enzyme important in health and disease. Physiol. Rev. 98, 641–665 (2018).
15. Munder, M. et al. Th1/Th2-regulated expression of arginase isoforms in murine macrophages and dendritic cells. J. Immunol. 167, 3771–3777 (1999).
16. Rath, M., Muller, I., Kopf, P., Cloos, E. I. & Munder, M. Metabolism via arginase or nitric oxide synthase: two competing arginine pathways in macrophages. Front Immunol. 5, 532 (2014).
17. Greenhalgh, A. D. et al. Arginase-1 is expressed exclusively by infiltrating myeloid cells in CNS injury and disease. Brain Behav. Immun. 56, 61–67 (2016).
18. Zarruk, J. G., Greenhalgh, A. D. & David, S. Microglia and macrophages differ in their inflammatory profile after permanent brain ischemia. Exp. Neurol. 301(Pt B), 120–132 (2018).
19. Shoshita, E. et al. Arginase 2 promotes neurovascular degeneration during ischemia/reperfusion injury. Cell Death Dis. 7, e2483 (2016).
20. Yau, T. et al. Preliminary efficacy, safety, pharmacokinetics, pharmacodynamics and quality of life study of pegylation recombinant human arginase 1 in patients with advanced hepatocellular carcinoma. Invest. New Drugs 33, 496–504 (2015).
21. Morrow, K. et al. Anti-lemuric mechanisms of pegylated arginase I in acute lymphoblastic t-cell leukemia. Leukemia 27, 569–577 (2013).
22. Musai, F. et al. Arginina dependence of acute myeloid leukemia blast proliferation: a novel therapeutic target. Blood 125, 2386–2396 (2015).
23. Fletcher, M. et al. L-arginine depletion blunts antitumor T-cell responses by inducing myeloid-derived suppressor cells. Cancer Res. 75, 275–285 (2015).
24. Feun, L. G., Kuo, M. T. & Sava, N. Arginase deprivation in cancer therapy. Curr. Opin. Clin. Nutr. Metab. Care 18, 78–82 (2015).
25. Jyer, R. K. et al. Mouse model for human arginase deficiency. Mol. Cell Biol. 22, 4491–4498 (2002).
26. Yokota, H. et al. Neuroprotection from retinal ischemia/reperfusion injury by NOX2 NADPH oxidase deletion. Invest. Ophthalmol. Vis. Sci. 52, 8123–8131 (2011).

27. Zheng, L., Gong, B., Hatala, D. A. & Kern, T. S. Retinal ischemia and reperfusion causes capillary degeneration: similarities to diabetes. Invest. Ophthalmol. Vis. Sci. 48, 361–367 (2007).

28. Rosenbaum, D. M. et al. The role of the p53 protein in the selective vulnerability of the inner retina to transient ischemia. Invest. Ophthalmol. Vis. Sci. 39, 2132–2139 (1998).

29. Schmid, H., Renner, M., Dick, H. B. & Joachim, S. C. Loss of inner retinal neurons after retinal ischemia in rats. Invest. Ophthalmol. Vis. Sci. 55, 2777–2787 (2014).

30. Kim, C. R., Kim, J. H., Park, H. L. & Park, C. K. Ischemia reperfusion injury triggers TUNA-induced necroptosis in rat retina. Curr. Eye Res. 42, 771–779 (2017).

31. Gao, S., Andreea, K. & Cooper, N. G. Ischemia-reperfusion injury of the retina is linked to necroptosis via the ERK1/2-AMPK pathway. Mol. Vis. 20, 1347–1387 (2014).

32. Divianitchkova, G., Degterev, A. & Ivanov, D. Retinal ganglion cell (RGC) programmed necrosis contributes to ischemia-reperfusion-induced retinal damage. Exp. Eye Res. 123, 1–7 (2014).

33. Rosenbaum, D. M. et al. Necroptosis, a novel form of caspase-independent cell death, contributes to neuronal damage in a retinal ischemia-reperfusion injury model. J. Neurosci. Res. 88, 1569–1576 (2010).

34. Park, S. W. et al. A selective inhibitor of DRP1, Mdiv-1, increases retinal ganglion cell survival in acute ischemic mouse retina. Invest. Ophthalmol. Vis. Sci. 52, 2893–2894 (2011).

35. Vininguniarpetj, I. A. et al. NLRP3 inflammasome activation drives bystander cone photoreceptor cell death in a P23H rhodopsin model of retinal degeneration. Hum. Mol. Genet. 25, 1501–1516 (2016).

36. Ahmed, A. et al. Minocycline modulates microglia polarization in ischemia-reperfusion model of retinal degeneration and induces neuroprotection. Sci. Rep. 7, 14067 (2017).

37. Van den Bossche, J., Baardman, J. & de Winther, M. P. Metabolic character-