



VASCULAR BIOLOGY, ATHEROSCLEROSIS, AND ENDOTHELIUM BIOLOGY

The Inability of the Choroid to Revascularize in Oxygen-Induced Retinopathy Results from Increased p53/miR-Let-7b Activity



Tianwei E. Zhou,^{*†} Tang Zhu,[‡] José C. Rivera,^{†‡§} Samy Omri,[†] Houda Tahiri,[†] Isabelle Lahaie,[†] Raphaël Rouget,[‡] Maëlle Wirth,[†] Stanley Nattel,^{*¶} Gregory Lodygensky,^{||} Gerardo Ferbeyre,^{**} Mohammad Nezhady,^{††} Michel Desjarlais,^{*†} Patrick Hamel,[§] and Sylvain Chemtob^{*†‡§||}

From the Department of Pharmacology and Therapeutics,* McGill University, Montréal, Québec; and the Department of Ophthalmology,[†] Maisonneuve-Rosemont Hospital Research Centre, the Departments of Pharmacology,[‡] Ophthalmology,[§] and Pediatrics,^{||} Sainte-Justine University Hospital Centre, the Department of Medicine,[¶] Montreal Heart Institute, and the Departments of Biochemistry** and Pathology and Cell Biology,^{††} University of Montréal, Montréal, Québec, Canada

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Address correspondence to
Sylvain Chemtob, M.D., Ph.D.,
or Tianwei E. Zhou, M.D.,
Ph.D., Department of Ophthalmology,
Maisonneuve-Rosemont Hospital Research
Centre, University of Montreal,
5415 Blvd de l'Assomption,
Montreal, QC H1T 2M4, Canada.
E-mail: sylvain.chemtob@umontreal.ca or ellen.zhou@umontreal.ca.

Retinopathy of prematurity (ROP) is characterized by an initial retinal avascularization, followed by pathologic neovascularization. Recently, choroidal thinning has also been detected in children formerly diagnosed with ROP; a similar sustained choroidal thinning is observed in ROP models. But the mechanism underlying the lack of choroidal revascularization remains unclear and was investigated in an oxygen-induced retinopathy (OIR) model. In OIR, evidence of senescence was detected, preceded by oxidative stress in the choroid and the retinal pigment epithelium. This was associated with a global reduction of proangiogenic factors, including insulin-like growth factor 1 receptor (Igf1R). Coincidentally, tumor suppressor p53 was highly expressed in the OIR retinae. Curtailing p53 activity resulted in reversal of senescence, normalization of Igf1r expression, and preservation of choroidal integrity. OIR-induced downregulation of Igf1r was mediated at least partly by miR-let-7b as i) let-7b expression was augmented throughout and beyond the period of oxygen exposure, ii) let-7b directly targeted *Igf1r* mRNA, and iii) p53 knock-down blunted let-7b expression, restored Igf1r expression, and elicited choroidal revascularization. Finally, restoration of Igf1r expression rescued choroid thickness. Altogether, this study uncovers a significant mechanism for defective choroidal revascularization in OIR, revealing a new role for p53/let-7b/IGF-1R axis in the retina. Future investigations on this (and connected) pathway could further our understanding of other degenerative choroidopathies, such as geographic atrophy. (*Am J Pathol* 2019, 189: 2340–2356; <https://doi.org/10.1016/j.ajpath.2019.07.009>)

Retinopathy of prematurity (ROP) is the leading cause of severe visual impairment and blindness in neonates and young children in North America and Europe.¹ ROP has long been associated with damage of the inner retina, including dysregulated angiogenesis of retinal vessels.² Recently, clinical and preclinical animal studies have detected choroidal thinning in subjects previously afflicted with ROP^{3–6} (Supplemental Figure S1), which can result in photoreceptor injury.^{7,8} But unlike retinal vessels that undergo neovascularization on retinal ischemia and hypoxia, the choroid—the exclusive source of oxygen and nutrients to photoreceptors⁹—does not revascularize and remains thin

despite ensuing local hypoxia.⁷ The underlying mechanisms for such paradoxical persistent choroidal thinning remain unexplained.

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Along these lines, early senescence is a process that promotes age-related disorders; and when applied to cell replication, it interferes with proliferation. In this context, oxidative stress and hyperoxia, as well as hypoxia and nutrient deficiency to the retinal pigment epithelium (RPE) and photoreceptors secondary to choroidal involution,⁷ are established triggers of senescence.¹⁰ Oxygen supplementation and deficient antioxidant defense in premature subjects exert a major contribution in the development of ROP/oxygen-induced retinopathy (OIR).² Ensued increased generation of reactive oxygen species from the oxidative stress readily induces DNA damage¹⁰ and subsequently activates ataxia-telangiectasia mutated serine/threonine kinase—a potent activator of p53.¹¹ Hypoxic ischemia, at times linked to oxidative stress, elicits an endoplasmic reticulum stress and unfolded protein response to limit protein translation, which concomitantly increases degradation of mRNA through activation of inositol-requiring enzyme 1 α endoribonuclease, followed by induction of p53^{12,13} contributing to the senescent phenotype.¹⁴ Through its properties as a transcription factor and others,¹⁵ p53 tightly regulates cell fate, including cell cycle progression, apoptosis, and senescence—all aiming to maintain genomic stability and avoid passage of DNA damages to daughter cells.¹⁶ In particular, p53 governs replicative cellular senescence by targeting p21/cyclin-dependent kinase inhibitor 1 and different miRNAs.¹⁷

The expression of p53 is largely regulated by miRNAs¹⁸ and, in turn, p53 controls the expression of various miRNAs involved in cell fate and angiogenesis.¹⁹ miRNAs are a family of noncoding RNAs that fine-tune post-transcriptional regulation of genes. On the basis of knowledge mostly acquired from cancer research, p53 regulates angiogenic pathways via certain families of miRNAs—for instance, the p53/miR-200 pair targets the vascular endothelial growth factor (VEGF) pathway,^{20,21} whereas p53/let-7 modulates the critical insulin-like growth factor 1 (IGF1) pathway at multiple levels.^{22–24} But in retinal diseases, mechanistic insights about the interactions of p53, miRNAs, and their downstream pathways are scarce. Herein, we propose to investigate the unexplored role of p53 and regulatory miRNAs in contributing to deficient choroidal revascularization in the context of ROP. This study also explores the plausible link between cellular stress experienced in the neonatal period with early senescence.

Materials and Methods

Animals

Pigmented Long-Evans rats were purchased from Charles Rivers Laboratories (Saint-Constant, Canada). All experiments were conducted in accordance with the Association for Research in Vision and Ophthalmology statement regarding use of animals in ophthalmic and vision research

and were approved by the Hôpital Maisonneuve—Rosemont Animal Care Committee.

Using Optical Coherence Tomography to Measure Choroidal Thickness

A portable, noncontact, hand-held, high-resolution optical coherence tomography instrument (Envisu C2300; Leica Microsystems, Wetzlar, Germany) was used. Patients were imaged in the neonatal intensive care unit at Sainte-Justine Hospital during ROP screening. Informed consent for publication was obtained from parents or legal guardians.

Patients were imaged without the use of anesthetic or eyelid speculum. Choroidal thickness, from the posterior edge of the RPE to the choroid/sclera junction, was measured with the built-in caliper, applied perpendicularly to the RPE, just under the fovea.

OIR Model

Newborn Long-Evans rats were placed with their mothers under oxygen concentration, which cycles at 50% \pm 1% or 10% \pm 1% every 24 hours from postnatal day (P) 0 to P14. On P14, all rats were returned to room air. This model is characterized by a retinal avascularization predominantly in the periphery (at P14), followed by hypoxia-driven retinal neovascularization clearly present at P18, which mimics the two principal phases of ROP in humans.^{25,26} Controls were maintained in room air (21% O₂). In some cases, pups were selected to receive twice-daily i.p. injections of p53 inhibitor [pifithrin- α (PFT α); 20 mg/kg; Sigma-Aldrich, St. Louis, MO] from P0 to P14. Animals were sacrificed at different ages (P3, P6, P10, P14, P18, and P30), and eyes were collected and processed for measurements described below.

Separation of RPE Cells and Choroid

Pigmented Long-Evans rats facilitated dissection of RPE cells from choroid. Rats were sacrificed at various ages (P3 to P30, as described above), and eyes were enucleated. Eyes were washed in icy phosphate-buffered saline (catalog number 311-012-CL; Wisent Inc., Saint-Jean-Baptiste, Canada) with gentle shaking for 10 minutes to remove any attached animal hair. The cornea and lens were subsequently removed. The freshly isolated eye cups, which contained neuroretina, RPE, choroid, and sclera, were incubated in Hanks' balanced salt solution (catalog number 14025076; Thermo Fisher Scientific, Waltham, MA) with 2.5% TrypLE Express Enzyme (catalog number 126040132; Thermo Fisher Scientific) and 2 mg/mL Collagenase D (catalog number 11088866001; Roche Life Science, Branford, CT) at 37°C for 45 minutes. Eye cups were washed using Dulbecco's modified Eagle's medium (DMEM)/Nutrient Mixture F-12 (catalog number 319-075-CL; Wisent Inc.) to finalize the digestion step and then kept

on ice. The neuroretina immersed in DMEM/Nutrient Mixture F-12 was separated from RPE/choroid complex using a pair of blunt-headed forceps to gently tap on the choroid and dislodge RPE cells; then, RPE cells were recovered using a pipette. RPE cells from four eyes were pooled and centrifuged. DMEM/Nutrient Mixture F-12 solution was removed and RiboZol (catalog number N580-200ML; Amresco, Solon, OH) was added to RPE cells for RNA isolation (see below).

RNA Isolation and Real-Time Quantitative PCR

Eyes were rapidly enucleated and placed into a sterile Petri dish resting on ice. The RPE/choroid complex was promptly separated from neuroretina. Both tissue complexes were processed for RNA using RiboZol, followed by treatment with DNase I (catalog number AMPD1; Sigma-Aldrich) to remove any contaminating genomic DNA. The DNase-treated RNA was then converted into cDNA using iScript Reverse Transcription Kit (catalog number 170-8841; Bio-Rad, Mississauga, Canada). PCR primers targeting rat were designed using National Center for Biotechnology Information Primer Blast; detailed sequences are summarized in [Table 1](#). Quantitative analysis of gene expression was generated using an ABI 7500 Real-Time PCR (Applied Biosystems, Foster City, CA) sequence detection system and the SYBR Green Master Mix Kit (catalog number 1725271; BioRad). Gene expression was calculated relative to 18S universal primer pair (catalog number AM1718;

Ambion, Waltham, MA) expression using the ΔC_T method. Data were quantified by $\Delta\Delta C_T$ method.

PCR Array

RNA was isolated from RPE/choroid complex using the RiboZol method described in [Separation of RPE Cells and Choroid](#) and [RNA Isolation and Real-Time Quantitative PCR](#). RNA was then converted into cDNA using iScript Reverse Transcription Kit (catalog number 170-8841; Bio-Rad). Real-time PCR for angiogenic gene profiling was performed using the SYBR Green–Based Rat Angiogenesis PCR Array (catalog number PARN-024Z; Qiagen, Hilden, Germany).

miRNA PCR Array

RNA contents were isolated using miRNeasy Mini Kit (catalog number 217004; Qiagen), as per manufacturer's instruction. Total RNA (250 ng) was reverse transcribed with the miScript II RT kit (catalog number 218161; Qiagen), according to manufacturer's guidelines. cDNA (20 μ L) was diluted with RNase-free water before PCR. Real-time PCR for mature miRNA expression profiling was developed using the SYBR Green–based rat hypoxia signaling pathway miScript miRNA PCR array (catalog number MIRN-121ZA; Qiagen); miRNA primers are shown in [Table 2](#). The data obtained were analyzed using the $\Delta\Delta C_T$ method of relative quantification for miScript miRNA PCR arrays with Qiagen online software (<https://www.qiagen.com/at/shop/genes-and-pathways/data-analysis-center-overview-page>). Fold changes in miRNA between normoxia- and OIR-raised rats, with $P < 0.05$, were considered statistically significant; miRNAs with statistically significant fold changes were represented separately.

Microvascular Sprouting from Choroid Explants

Choroid from juvenile rats was prepared, as previously described.²⁷ Briefly, choroid was cut into approximately 2- × 1-mm pieces and placed in Matrigel (catalog number 354230; Corning, Corning, NY) with 2% fetal bovine serum and incubated in 37°C, 5% CO₂ for 48 hours before treatment. Choroidal tissue was obtained from control (normoxia) and OIR animals. For each type of *in vivo* condition, the left eye was cultured in normoxia for 48 hours

Table 1 qPCR Primer List

Gene symbols	Sequences
<i>Cdkn1a</i>	F: 5'-AGTATGCCGTCGTCTGTTTCG-3' R: 5'-GAGTGCAAGACAGCGACAAG-3'
<i>Tp53</i>	F: 5'-ACAGCGTGGTGGTACCGTAT-3' R: 5'-GGAGCTGTTGCACATGTA-3'
<i>Nfe2l2</i>	F: 5'-TTGTAGATGACCATGAGTCGC-3' R: 5'-CTCCATGTCTGCTGTATGC-3'
<i>Sod1</i>	F: 5'-CGGATGAAGAGGGCATGTT-3' R: 5'-CAATCACACCACAAGCCAAG-3'
<i>Sod2</i>	F: 5'-GCGACCTACGTGAACAATCT-3' R: 5'-CAGCAACTCTCCTTTGGGTT-3'
<i>Nox1</i>	F: 5'-CCCTTTGCTTCTTCTTGAATC-3' R: 5'-GCACCCGTCTCTTACAAATCC-3'
<i>Nox2</i>	F: 5'-TGATCATCACATCCTCCACCA-3' R: 5'-GATGGCAAGGCCGATGAA-3'
<i>Nox3</i>	F: 5'-GCAGCATTGGCGTGTCTT-3' R: 5'-GAAATGAACGCCCTAGGATCT-3'
<i>Nox4</i>	F: 5'-CTGCATCTGCTGAACTCAA-3' R: 5'-TCTCTGCTAGGGACCTTCTGT-3'
<i>Igf1r</i>	F: 5'-TGGCAGAAGTCTGTCTGAG-3' R: 5'-AACGCAGGGTCTAGTTGAGC-3'
<i>Igf1</i>	F: 5'-GCATTGTGGATGAGTGTTC-3' R: 5'-GGTCTTGTCTTCTGCACTTC-3'

F, forward; qPCR, real-time quantitative PCR; R, reverse.

Table 2 miRNA qPCR Primer List

Official symbol	Gene symbols
miRlet7b	MS00000007
miRlet7d	MS00012915
miR200a	MS000000581
miR203	MS00000602
miR29b-1	MS00005544

qPCR, real-time quantitative PCR.

in EGM-2MV medium (catalog number CC-4147; Lonza, Walkersville, MD), and the right eye was first placed in hypoxia (5% O₂) for 24 hours and then returned to normoxia for an additional 48 hours (also in EGM-2MV medium); in each case, tissues were cultured in presence or absence of 25 μmol/L p53 inhibitor PFTα.

Images of individual explants were taken before and after treatments. The vascular sprouting covered areas were quantified with ImageJ bundled with Java version 1.8.0_172 (<https://imagej.nih.gov/ij/download.html>); developed by Wayne Rasband, NIH, Bethesda, MD).

Ocular Tissue Preparation for Immunohistochemistry

Animals were perfused with phosphate-buffered saline and 4% paraformaldehyde (catalog number P6148; Sigma-Aldrich). For histology sections, eyes were immediately collected, dehydrated with alcohol, and embedded in paraffin. Sagittal sections (5 μm thick) were cut using a microtome (RM 2145; Leica Biosystems, Wetzlar, Germany). Eyes for cryopreparation were further fixed in 4% paraformaldehyde overnight. Posterior eye cups were frozen in OCT medium (catalog number 3801480; Leica Biosystems). Samples were then cut into sagittal sections (10 μm thick; CM3050S; Leica Biosystems) and processed for immunohistochemistry.

Choroidal Thickness Measurement

To measure choroidal thickness, cryosection samples of retina/choroid were stained with tetramethylrhodamine-conjugated lectin endothelial cell marker (dilution 1:200; Sigma-Aldrich; L5264). Choroidal images were then captured by confocal microscope (Olympus, Richmond Hill, Canada). Starting from the optic nerve (the 0 μm position), choroidal thickness was measured at 150-μm intervals across the choroid; measurements were averaged and plotted into a spider graph. Because the rat eye changes size from P14 to P210, the thickness was measured at one-quarter of the entire choroidal length, starting from the optic nerve, which corresponds to the region most affected by OIR.^{3–6}

Western Blot Analysis

Protein extracts (40 μg) were loaded, separated by voltage potential, and transferred to polyscreen polyvinylidene difluoride transfer membrane (catalog number 1620177; BioRad), as previously reported.²⁸ Respective proteins were revealed by antibodies listed in Table 3.

DNA Fragmentation Labeling

Labeling of fragmented DNA was performed by using a commercial kit based on terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling technique

Table 3 Antibody List

Protein	Company (catalog number)	Dilution used
β-Actin	Novus Biologicals, Oakville, Canada (NB600-501)	1:4000
IGF-IRβ	Santa Cruz Biotechnology, Dallas, TX (SC-713)	1:100
Phos-IGF-IRβ (Tyr 1161)	Santa Cruz Biotechnology (SC-101703)	1:100
p53	Santa Cruz Biotechnology (SC-393031)	1:100

IGF-IRβ, insulin-like growth factor 1 receptor β; Phos-IGF-IRβ, phosphorylated IGF-IRβ.

(catalog number 11684809910; Roche Life Science, Branford, CT).

siRNA Preparation

siRNAs were synthesized using an siRNA Construction Kit (catalog number AM1620; Ambion). The target sequences of the human p53-siRNA were designed by an siRNA design tool (Chang Bioscience, Fremont, CA) based on the human p53 DNA sequence (<https://www.ncbi.nlm.nih.gov/nucore>; accession number NM_000546.5).

Three oligonucleotides of human p53 were selected as the target area, which were as follows: p53-siRNA70 [sense, 5'-GAAACTTCTTCCTCCAGATGA-3' (forward)], p53-siRNA741 [sense, 5'-GGCCCATCCTTACCATCATCA-3' (forward)], and p53-siRNA969 [sense, 5'-GAGAA-TATTTACCCCTTAAGA-3' (forward)]. For the siRNA hairpin loop construction, AA and CCTGTCTC codes were added into the 5'- and 3'-terminal of the selected oligonucleotides, respectively. The designed hairpin oligonucleotides were then synthesized by Alpha-DNA Inc. (Montreal, Canada).

The synthesized siRNA hairpin oligonucleotides were used for the siRNA synthesis on the instruction sheet of the siRNA Construction Kit (catalog number AM1620; Ambion), which underwent the processes of transcription template preparation, double-stranded RNA synthesis, and siRNA preparation and purification.

To characterize the siRNAs, a rat retinal ganglion cell line was used (retinal ganglion cell line was kindly provided by Dr. Neeraj Agarwal, National Eye Institute, Bethesda, MD). Retinal ganglion cells were transfected with siRNA70, siRNA741, and siRNA969, as previously reported^{29,30}; and real-time quantitative PCR was performed to confirm their knock-down efficiencies.

In Silico Prediction of Potential miRNA Targets

miRNAs that can potentially target rat Igf1 receptor (Igf1r) transcript were predicted via TargetScan version 6.2 (TargetScan, Cambridge, MA) by searching for the presence of

well-conserved 8-mer (2636 to 2643) and 7-mer (6674 to 6680) sites on the 3'-untranslated region (3'UTR) of *Igf1r* mRNA. PicTar (<http://pictar.mdc-berlin.de>; last accessed February 7, 2019) and DIANA-microT-CDS (<http://www.microna.gr/microT-CDS>; last accessed February 7, 2019) were used to confirm TargetScan findings.

Construction of Luciferase-miRNA-Target Site Fusion Plasmids

The pmirGLO Dual-Luciferase miRNA Target Expression Vector (catalog number E1330; Promega, Madison, WI) was provided by S.N. To construct reporter vectors bearing miRNA-target sites, fragments containing the exact target sites for let-7b in the 3'UTR of rat *Igf1r* mRNA were synthesized (Top Gene Technologies Inc., Montreal, Canada). Each construct contained one of the two predicted binding sites for let-7b. These inserts were ligated into PmeI and XbaI sites on the pmirGLO Dual-Luciferase miRNA Target Expression Vector.

The Dual-Luciferase Reporter Assay

The dual-luciferase assay was performed with Dual-Luciferase Reporter Assay System (catalog number E1910; Promega). ARPE-19 cells (catalog number CRL-2302; ATCC, Manassas, VA) or rat brain microvascular endothelial cells (catalog number R840-05a; Cell Applications Inc., San Diego, CA) were seeded onto a 96-well plate at 2×10^4 cells/well. On the following day, 0.337 μg of the reporter vector (eg, *Igf1r*-3'UTR-pmirGLO-1/2) with miRNA mimic Syn-rno-let-7b-5p (catalog number MSY0000775; Qiagen) (working concentration = 20 nmol/L) was transfected into ARPE-19 cells using 1.684 μL of polyethylenimine transfection reagent (Polysciences, Warrington, PA) at 1 mg/mL. AllStars Negative Control siRNA (catalog number SI03650318; Qiagen) was used as the negative control. The luciferase reporter assay was performed 48 hours after transfection. Standard protocol, provided by the manufacturer, was followed. Briefly, ARPE-19/rat brain microvascular endothelial cells were lysed with Passive Lysis Buffer (20 μL /well) and gently shook at room temperature for 15 minutes. The assay plate was then inserted into an EnVision Microplate screening fluorescence reader (catalog number 2104-0010A; PerkinElmer, Waltham, MA). As soon as 75 μL Luciferase Assay Reagent II (Promega) was injected, the firefly luciferase signal was measured at 640 nm. Next, 75 μL of the Stop & Glo Reagent Buffer (Promega) was injected and the Renilla luciferase signal (used as an internal control) was measured at 525 nm. The firefly OD/Renilla OD ratio was calculated. The activity of Syn-has/rno-let-7b-5p was calculated by subtracting the firefly OD/Renilla OD ratio of samples that received AllStars Negative Control miRNA from that of samples that received Syn-has/rno-let-7b.

Preparation of Rat IGF1R Lentivirus for Subretinal Injection

The basic protocol of the lentiviral plasmid transfection has been previously described³¹ and modified. To prepare lentiviral particles, 9 μg of rat *Igf1r* (<https://www.ncbi.nlm.nih.gov/nucleotide>; accession number NM_052807.2; catalog number LVP933; Genetarget, San Diego, CA) plasmid was mixed with 6 μg of pRSV-Rev (Addgene, Cambridge, MA), 7.8 μg of pCMV-vesicular stomatitis virus glycoprotein (Addgene), and 15 μg of pMDLg/pRRE plasmid (Addgene), in 1.75 μL DMEM (catalog number 11995065; Thermo Fisher Scientific). Polyethylenimine linear (140 μL of 1 mg/mL; catalog number 23966; Polysciences Inc., Warrington, PA) in 1.61 mL of DMEM was added to the *Igf1r* plasmid mixture. Fifteen minutes later, the mixed transfection reagent was pipetted into a 15-cm-diameter culture dish preseeded with HEK293T cells at 50% to 60% confluence in 15 mL of complete DMEM. In total, four dishes were applied for lentiviral particle concentration. Twenty-four hours later, the transfecting medium was replaced with 18 mL complete DMEM. After a 30-hour incubation, the maximal secreted lentiviral particles in the medium were collected and sterilized by passing through a 0.22- μm filter (catalog number GSWP01300; Merck Millipore Ltd, Co, Cork, Ireland). For subretinal injection, the fresh lentiviral particle medium was concentrated by ultracentrifugation at $50,000 \times g$ for 4 hours. The supernatant medium was carefully removed, and the remaining pellet (concentrated lentivirus particle) was resuspended in 30 μL of sterilized phosphate-buffered saline by shaking ($8 \times g$) at 4°C for overnight. The final viral particles were gently mixed by pipetting, aliquoted into 10 μL /each into PCR microtubes, and stocked at -80°C until use. The concentration of the purified lentiviral particles was detected by Lenti-X p24 Rapid Titer Kit (catalog number 632200; Takara, Kusatsu, Japan) on the kit menu instruction.

Subretinal Injections of siRNA or Lentivirus

Rat pups (P14) were anesthetized with 85 mg/kg ketamine (Vetalar; Bioniche Animal Health, Belleville, ON, Canada) and 5.0 mg/kg xylazine (Rompun; Bayer HealthCare, Mississauga, ON, Canada). Animals were placed at lateral recumbency under a dissecting microscope (SteREO Discovery V8; Zeiss, Toronto, Canada). The cornea was punctured slightly anterior to the ora serrata with a 23-gauge needle (catalog number 305143; Becton Dickinson, Franklin Lakes, NJ); the needle was advanced full thickness through the cornea into the anterior chamber. Extra care was taken to avoid injuring the lens. A slight resistance to the movement of the needle indicated penetration of the retina and entrance into the subretinal matrix; 1 μL of siRNA or lentivirus solution was injected into central subretinal regions because choroid

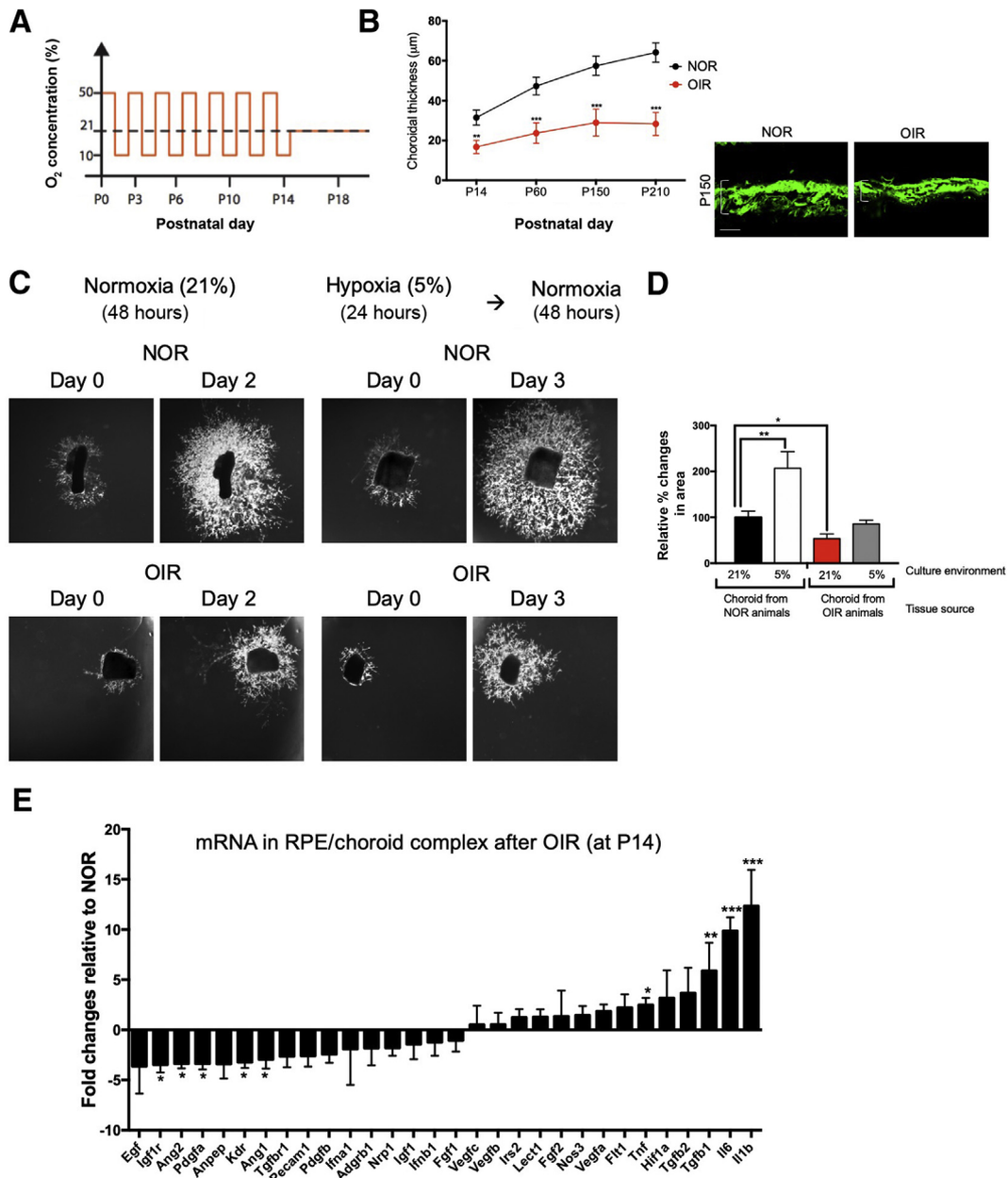


Figure 1 Choroid from oxygen-induced retinopathy (OIR)—subjected rats reveals diminished angiogenic capability. **A:** The cyclic OIR model was used in this study. Postnatal day (P) 1 rat pups were placed under alternating O₂ concentrations at 50% and 10%. The oxygen exposure period lasted 14 days, after which rats were returned to normoxia (NOR; indicated by the **black dashed line**). **B:** Choroidal involution in OIR animals persists beyond the oxygen exposure period into adult life. **Left panel:** Choroidal thicknesses of control and OIR rats were compared longitudinally. Choroids in the OIR group were significantly thinner (approximately 50%) than the NOR controls. **Right panel:** A representative immunohistochemistry image of choroidal thinning in OIR. The brackets outline choroidal thickness. *P* values are compared with corresponding NOR. **C:** *Ex vivo* choroidal explants were performed to assess angiogenic propensity. Choroidal tissues were obtained from rats under normoxia or OIR. One set of choroid tissues was cultured in normoxia for 48 hours; the other set was first placed in hypoxia (5% O₂) for 24 hours and returned to normoxia for another 48 hours. **D:** Under the normoxic conditions, choroidal explants from NOR rats expanded by 6.70-fold (black bar; set as a relative baseline value of 100% to which all others are compared). The sprouting area of OIR rats (red bar) only increased by 3.07-fold, merely 46% compared with NOR. Primed by hypoxia for 24 hours (followed by 48 hours in normoxia), choroids from NOR animals further expanded to 13.19-fold (197%; white bar); those from OIR choroid continued to expand for 4.86-fold (73%; gray bar). All changes shown in the histogram are relative to NOR cultured under normoxic conditions. *P* values are compared with the NOR choroid culture in normoxia (48 hours). **E:** Retinal pigment epithelium (RPE)/choroid complexes (from P14 rats) were analyzed by real-time quantitative PCR arrays for factors indicated; fold changes in expression are relative to NOR. A decrease in numerous proangiogenic factors in OIR, including vascular endothelial growth factor (Vegf) receptor 2 [alias kinase insert domain receptor (Kdr)], angiotensins 1 and 2 (Ang1/2), and most notably insulin-like growth factor 1 receptor (IGF1R; Igf1r), was detected. *P* values are compared with corresponding NOR. Data are expressed as means ± SEM (**B**, **D**, and **E**). *n* = 5 to 7 animals per group (**B**); *n* = 5 (**C** and **D**); *n* = 4 animals per group (**E**). **P* < 0.05, ***P* < 0.01, and ****P* < 0.001. Scale bar = 30 μm (**B**). Adgrb, adhesion G protein-coupled receptor B; Anpep, alanine aminopeptidase; Egf, epidermal growth factor; Fgf, fibroblast growth factor; Flt, Fms-like tyrosine kinase (alias Vegfr); Hif, hypoxia-inducible factor; Ifn, interferon; Irs, insulin receptor substrate; Lect, lectin; Nos, nitric oxide synthase; Nrp, neuropilin; Pecan, platelet endothelial cell adhesion molecule; Pdgf, platelet-derived growth factor; Tgfb, transforming growth factor β; Tgfb, TGF-β receptor; Tnf, tumor necrosis factor.

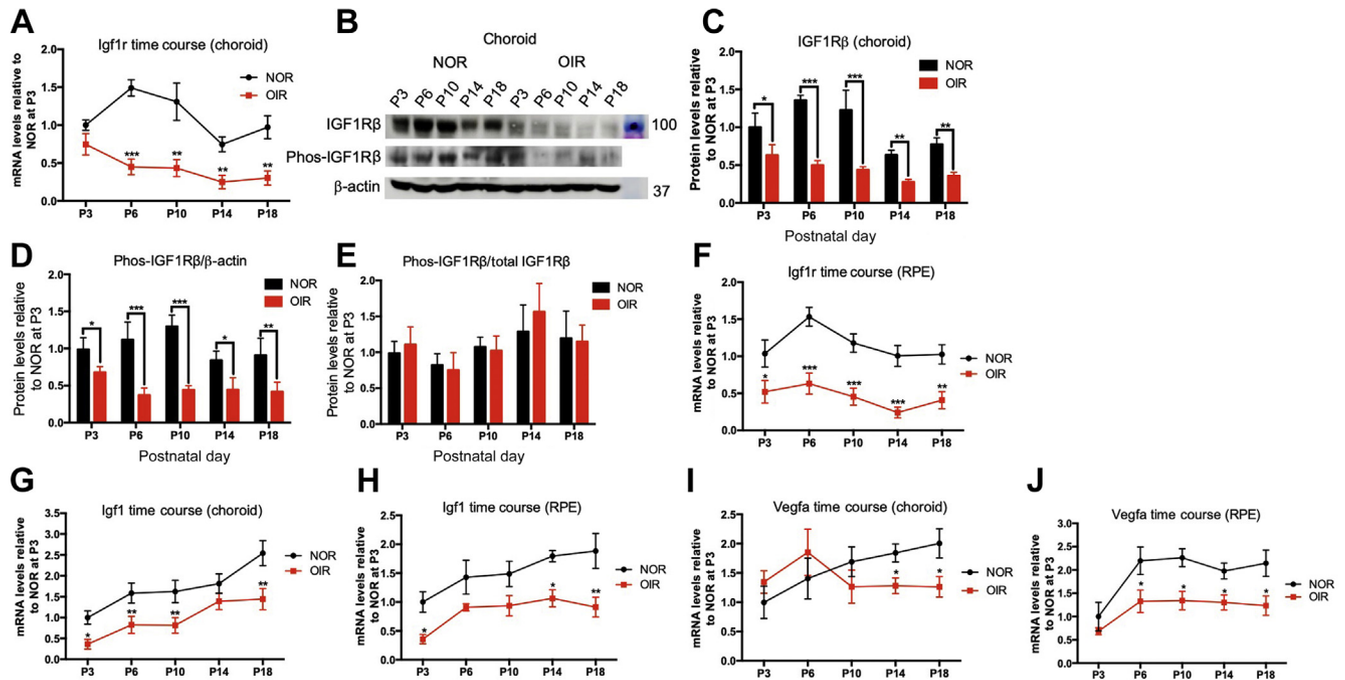


Figure 2 Down-regulation of insulin-like growth factor 1 (Igf1), Igf1 receptor (Igf1r), and vascular endothelial growth factor receptor 1 in retinal pigment epithelium (RPE) and choroid of oxygen-induced retinopathy (OIR)-subjected animals. **A:** Developmental changes in mRNA [by real-time quantitative PCR (qPCR)] and protein of Igf1r in choroid of normoxia (NOR)- and OIR-subjected rats. **B:** Representative Western blot analysis of Igf1r and phosphorylated (phos) Igf1r from postnatal day (P) 3 to P18. **C–E:** Western blot analyses presented in histogram: Igf1r protein (normalized by β -actin, indicating the amount of Igf1r; **C**); phos-Igf1r (normalized by β -actin, indicating the amount of phos-Igf1r; **D**); and phos-Igf1r/Igf1r (indicating the fraction of phosphorylated Igf1r; **E**) (please note that each ratio was further normalized to that of P3 NOR). **F:** Developmental changes in mRNA (by qPCR) of *Igf1r* in the RPE of NOR and OIR rats. **G** and **H:** Developmental changes in mRNA of *Igf1* in choroid and RPE of NOR- and OIR-subjected rats. **I** and **J:** Developmental changes in mRNA of *Vegfa* in choroid and RPE of NOR- and OIR-subjected rats. Data are expressed as means \pm SEM fold change from NOR set at 1 on P3 (**A** and **C–J**). $n = 4$ animals per group. * $P < 0.05$, ** $P < 0.01$, and *** $P < 0.001$ compared with corresponding NOR.

thinning mostly affects the central regions.^{3–6} The blunt end needle was then slowly retracted from the corneal opening to avoid reflux.

Statistical Analysis

Results are presented as means \pm SEM for all experiments. One- or two-way analysis of variance with significance $\alpha = 0.05$ was used for processing data. Bonferroni post hoc analysis was used to calculate significance between groups. Two-tailed t -tests were used to test for significance between two means.

Results

Persistent Choroidal Involution in OIR Animals Is Associated with Widespread Down-Regulation of Angiogenic Factors, including Igf1r

Using an established OIR model in rats by alternating exposure of pups to 50% and 10% O_2 every 24 hours from P0 to P14 (Figure 1A), choroidal thickness was measured at the end of the oxygen exposure and beyond. Reduced choroidal thickness was confirmed in OIR-subjected animals and was sustained into adult age (Figure 1B), despite

evidence for *in situ* hypoxia.⁷ Choroids from rats subjected to OIR exhibited limited ability to sprout in Matrigel, even on *ex vivo* exposure to hypoxia; hence, choroids from rats subjected to OIR displayed a reduced hypoxia-induced angiogenic propensity (Figure 1, C and D).

To determine possible mediators implicated in curtailing choroidal vascular repair during OIR, RPE/choroid complex isolated from OIR and control animals at P14 was assayed for an array of factors that affect angiogenesis (Figure 1E). PCR analysis showed a substantial increase in some inflammatory mediators, such as IL-6, tumor necrosis factor- α , and IL-1 β in RPE/choroid of OIR-exposed subjects, as previously reported,⁷ along with a widespread down-regulation of important vasosupportive factors, including angiopoietins 1 and 2, kinase insert domain receptor (alias Vegf receptor 2), platelet-derived growth factor A, and pronounced Igf1r (Figure 1E).

Because Igf1r is a prominent permissive angiogenic factor,³² a detailed age-dependent expression analysis of *Igf1r* and its ligand *Igf1* was conducted in choroid and RPE on exposure to supplemental oxygen. Experiments revealed sustained depression of both *Igf1r* and *Igf1* in OIR (Figure 2, A–H). In addition, the decrease in total Igf1r protein among OIR subjects was not compensated by activation (ie, phosphorylation) of the remaining Igf1r. The developmental profile of phosphorylated

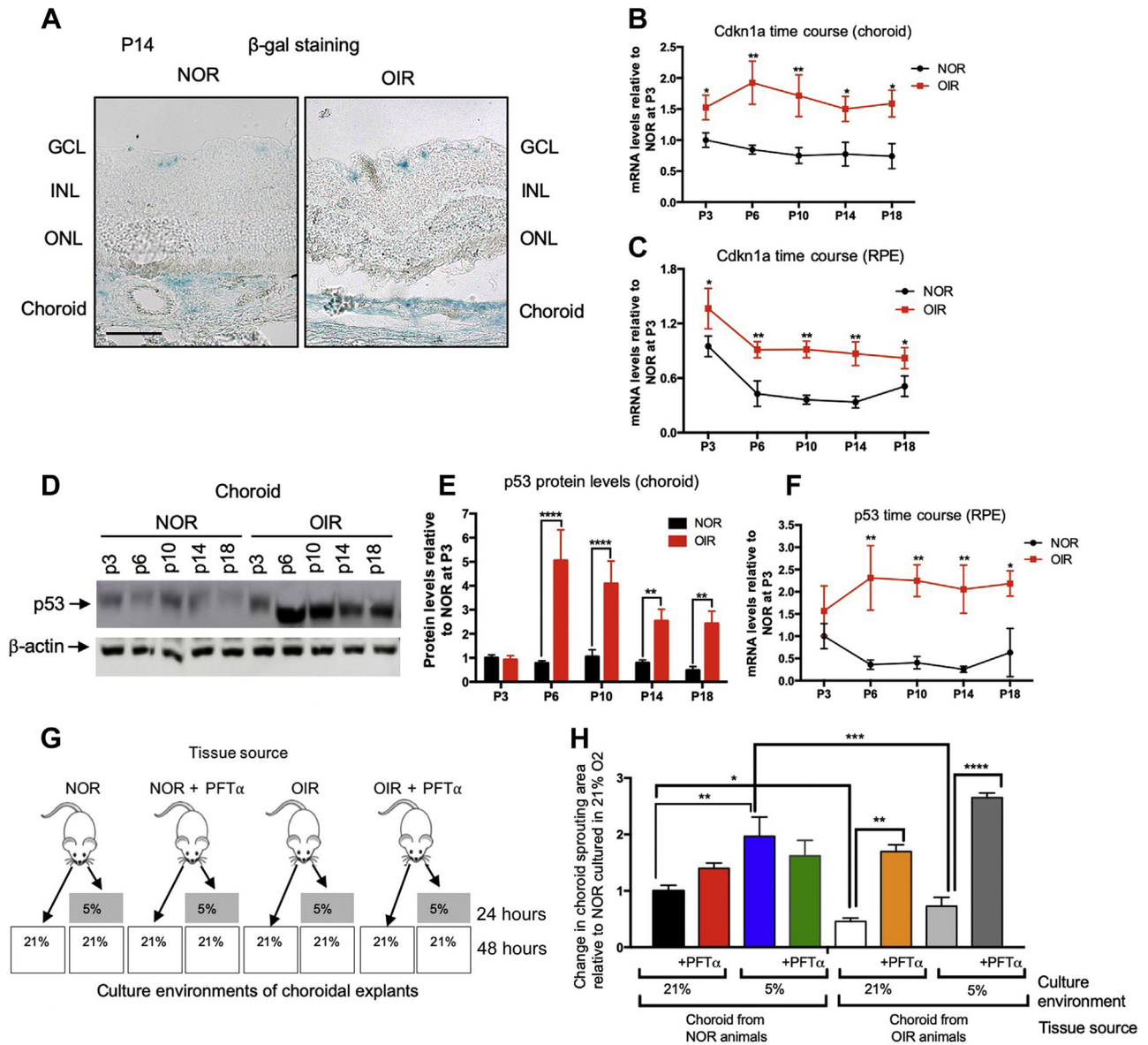


Figure 3 Senescence profile in choroid and retinal pigment epithelium (RPE) of normoxia (NOR)– and oxygen-induced retinopathy (OIR)–subjected rats. **A:** Representative senescence β -galactosidase (β -gal) staining shows robust subretinal signal in RPE and choroid of OIR-subjected rats at postnatal day (P) 14. **B** and **C:** Developmental profile of cyclin-dependent kinase inhibitor 1 (Cdkn1a; p21) mRNA in choroid and RPE of NOR- and OIR-subjected rats. **D–F:** Time course of p53 protein expression profile in choroid and mRNA in RPE of NOR- and OIR-subjected animals. **G:** *Ex vivo* choroidal explants were performed to assess vasoproliferative ability. Choroidal tissues were collected from NOR- and OIR-subjected rats treated with pifithrin- α (PFT α) or left untreated. One set of choroid tissues was cultured in normoxia for 48 hours, and another set was first placed in hypoxia (5% O $_2$) for 24 hours and returned to normoxia for an additional 48 hours. **H:** Changes in choroidal sprouting area are represented as relative to those of NOR-raised rats cultured in 21% O $_2$. Data are expressed as means \pm SEM (**B**, **C**, **E**, and **F**, fold change relative to NOR set at 1 on P3, and **H**, relative change). $n = 3$ independent samples (**A**); $n = 4$ animals per group (**B**, **C**, **E**, and **F**); $n = 5$ experiments (**H**); * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, and **** $P < 0.0001$ compared with corresponding NOR or as for indicated comparisons. Scale bar = 20 μ m (**A**). GCL, ganglion cell layer; INL, inner nuclear layer; ONL, outer nuclear layer.

Igf1r (Figure 2D) was parallel to that of total Igf1r (Figure 2C); the phosphorylated Igf1r/Igf1r ratio in the OIR group did not differ at different ages (Figure 2E). Together, these data suggested that the decrease in Igf1r activity in OIR rats is almost entirely due to reduced protein expression, rather than receptor activation. Because activation of Igf1r is a prerequisite for induction (among others) of Vegfa,^{33,34} Vegfa expression was

measured; it was significantly lower in choroid and RPE of OIR-subjected rats (Figure 2, I and J). Corresponding immunofluorescence microscopy analysis confirmed colocalization of Igf1r in choroidal endothelial cells, choroidal pericytes, and RPE cells from control animals, whereas Igf1r immunofluorescence was hardly detectable in samples from OIR-subjected rats (Supplemental Figure S2, A and B).

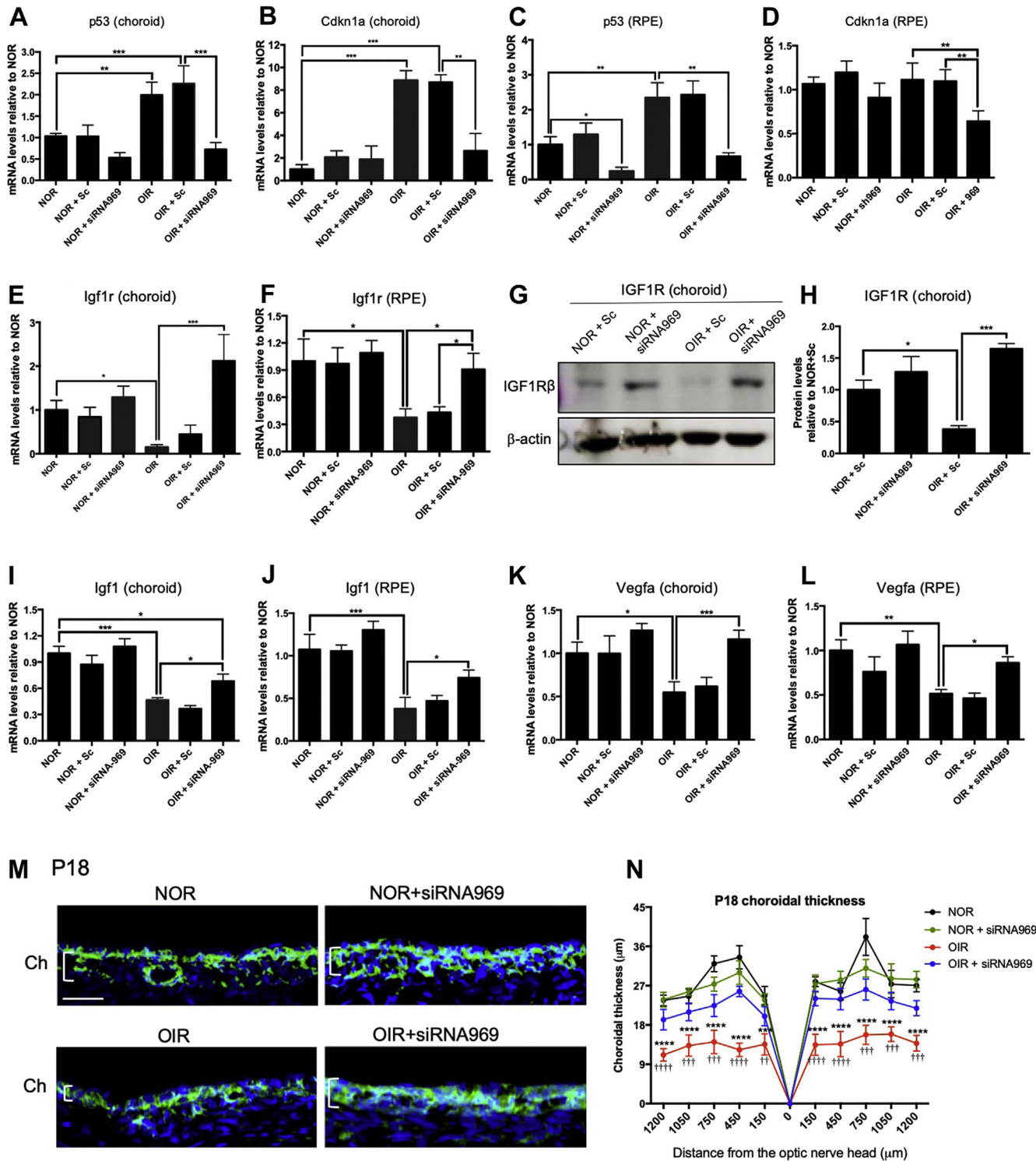


Figure 4 The p53 knockdown rescues insulin-like growth factor 1 receptor (Igf1r) [and vascular endothelial growth factor a (Vegfa)] expression and normalizes choroidal thickness. **A–F** and **I–L**: p53, cyclin-dependent kinase inhibitor 1 (Cdkn1a), Igf1r, Igf1, and Vegfa mRNA expression at postnatal day (P) 18 in central choroid and retinal pigment epithelium (RPE) of normoxia (NOR)— (set as arbitrary value of 1) and oxygen-induced retinopathy (OIR)—subjected animals treated with scrambled siRNA or (effective) siRNA969 that targets p53. Values fold change relative to NOR. **G** and **H**: Igf1r Western blot analysis and histogram quantification in choroid of NOR— (set as arbitrary value of 1) and OIR—subjected animals treated with scrambled siRNA or (effective) siRNA969. Values are fold change relative to NOR (set as 1). **M**: Representative choroid immunofluorescence of choroid stained with lectin (green; vasculature) and DAPI (blue; cell nuclei). The brackets outline choroidal thickness. **N**: Choroid thickness quantification in spider graph relative to distance from optic nerve (position 0). Values are given of choroid thickness (in μm). Data are expressed as means ± SEM (**A–F**, **H–L**, and **N**). *n* = 4 animals per group (**A–F** and **I–L**); *n* = 3 to 5 (**G** and **H**, experiments per group, and **N**, animals per group). **P* < 0.05, ***P* < 0.01, ****P* < 0.001, and *****P* < 0.0001 compared with corresponding NOR or as for indicated comparisons; ††*P* < 0.01, †††*P* < 0.001, and ††††*P* < 0.0001 compared with corresponding OIR rats treated with siRNA969. Scale bar = 30 μm (**M**). Ch, choroid; Sc, scrambled.

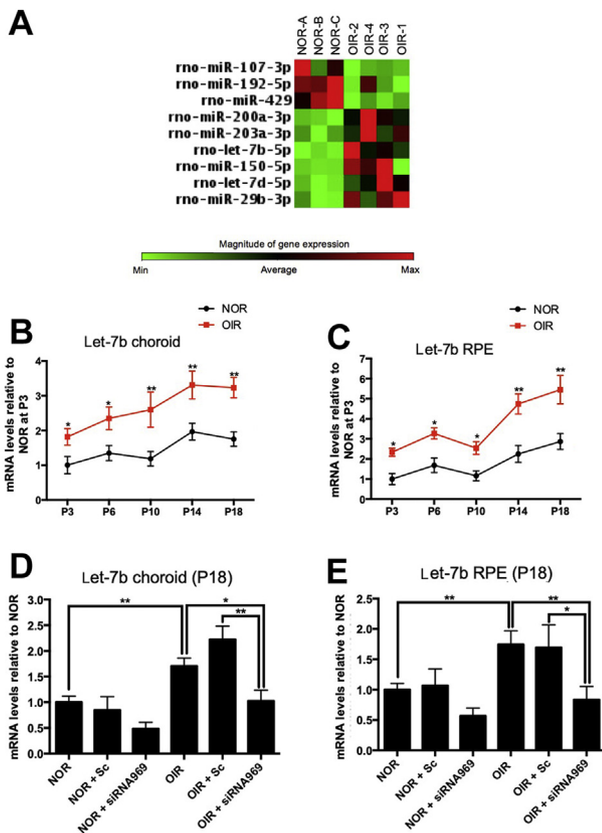


Figure 5 Relevant vasoproliferation-implicated miRNA profiling in choroid/retinal pigment epithelium (RPE) complex of animals subjected to oxygen-induced retinopathy (OIR). **A**: A heat map of most significantly altered miRNA expression in choroid/RPE complex of rats subjected to OIR relative to normoxia (NOR). Of 84 miRNAs that were assessed, only 8 showed statistically significant changes (increase or decrease). **B** and **C**: Time-course expression profile of let-7b in choroid and RPE of NOR- and OIR-subjected rats. Values are fold change from NOR (set as 1) on postnatal day (P) 3. **D** and **E**: Let-7b miRNA expression at P18 in choroid and RPE of NOR- (set as arbitrary value of 1) and OIR-subjected animals treated with scrambled siRNA or (effective) siRNA969 targeting p53. Values are fold change relative to NOR (set as 1). Data are expressed as means \pm SEM (**B–E**). $n = 3$ to 4 experiments (**A**); $n = 4$ animals per group (**B** and **C**); $n = 3$ to 6 experiments per group (**D** and **E**). * $P < 0.05$, ** $P < 0.01$ compared with corresponding NOR or as for indicated comparisons. Max, maximum; Min, minimum; rno, *Rattus norvegicus* (common rat); Sc, scrambled.

Evidence for Senescence and Up-Regulation of Tumor Suppressor p53 in Choroid and RPE of OIR-Subjected Animals

OIR is known to elicit oxidative stress,³⁵ which affects cell replication,^{36,37} through effects on a broad range of gene expression.³⁸ Increased oxidant stress in choroid and RPE in OIR witnessed soon (at P3) after oxygen exposure was confirmed. As anticipated, major reactive oxygen species generators *Nox1*, *Nox2*, and *Nox4* (but not significantly *Nox5*) were consistently up-regulated in choroid and RPE from rats subjected to OIR (Supplemental Figure S3, A–D and H–K); coherently, major antioxidant *SOD2* and, to a lesser extent, *SOD1* were down-regulated (Supplemental Figure S3, E, F, L, and M), and expression of major oxidative stress-activated

nuclear factor (erythroid-derived 2)-like 2 (*Nfe2l2*)³⁹ was augmented (Supplemental Figure S3, G and N).

Although hyperoxia and especially oxidative stress are important triggers of senescence,¹⁰ which can explain the inability of the choroid to revascularize (Figure 1, B–D), replicative cell senescence may also be contributed by local tissue hypoxia and nutrient deprivation, as seen in ischemia associated with choroidal thinning.⁷ Together with a decrease in growth factors, most prominently the critical Igf1r (Figure 1E), senescence-associated β -galactosidase and *Cdkn1a* (a prominent cell cycle arrest target) were assayed; and they were found to be robustly increased in choroid and RPE after OIR (Figure 3, A–C). These indicators of senescence were further supported by up-regulation of senescence-associated tumor suppressor p53, a pivotal regulator of cell fate (involving cell cycle, apoptosis, and senescence),^{40,41} which, of relevance, also controls transcription of *Cdkn1a*.⁴² More important, p53 expression was significantly elevated from P6 and beyond oxygen exposure in choroid and RPE of OIR-subjected animals (Figure 3, D–F).

Decreased Angiogenic Ability in Choroid of OIR-Subjected Animals Is Secondary to p53-Mediated Down-Regulation of Igf1r

To examine the antiangiogenic effect of p53, it was first shown that PFT α (a p53 inhibitor) abrogated OIR-induced increase in p53 and *Cdkn1a* in choroid and RPE (Supplemental Figure S4, A–C). This was associated with restoration of choroidal sprouting in response to hypoxia: PFT α -treated hypoxia-primed OIR choriocapillary grew more robustly than untreated tissue (Figure 3, G and H). Although p53 can also direct cell fate toward apoptosis,⁴³ there were no apparent apoptotic signals (notably, terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling-positive cells) in the choroid and RPE of OIR-subjected rats (Supplemental Figure S4D).

The antiangiogenic role of p53 was further ascertained by knocking down its expression. Of three siRNAs designed, siRNA969 was most effective in knocking down p53 *in vitro* (Supplemental Figure S5A); subretinal injection of siRNA969 (5 ng/mL; based on dose-response) was also

Table 4 miRNA Fold Changes

Mature identifier	Fold change	P value
rno-miR-192-5p	0.1863	0.0254
rno-miR-107-3p	0.2804	0.0444
rno-miR-429	0.3175	0.0019
rno-miR-150-5p	2.4114	0.0801
rno-miR-203a-3p	2.7856	0.0460
rno-let-7d-5p	3.2522	0.0318
rno-miR-29b-3p	3.3988	0.0185
rno-let-7b-5p	3.5211	0.0196
rno-miR-200a-3p	3.62	0.0151

rno, *Rattus norvegicus* (common rat).

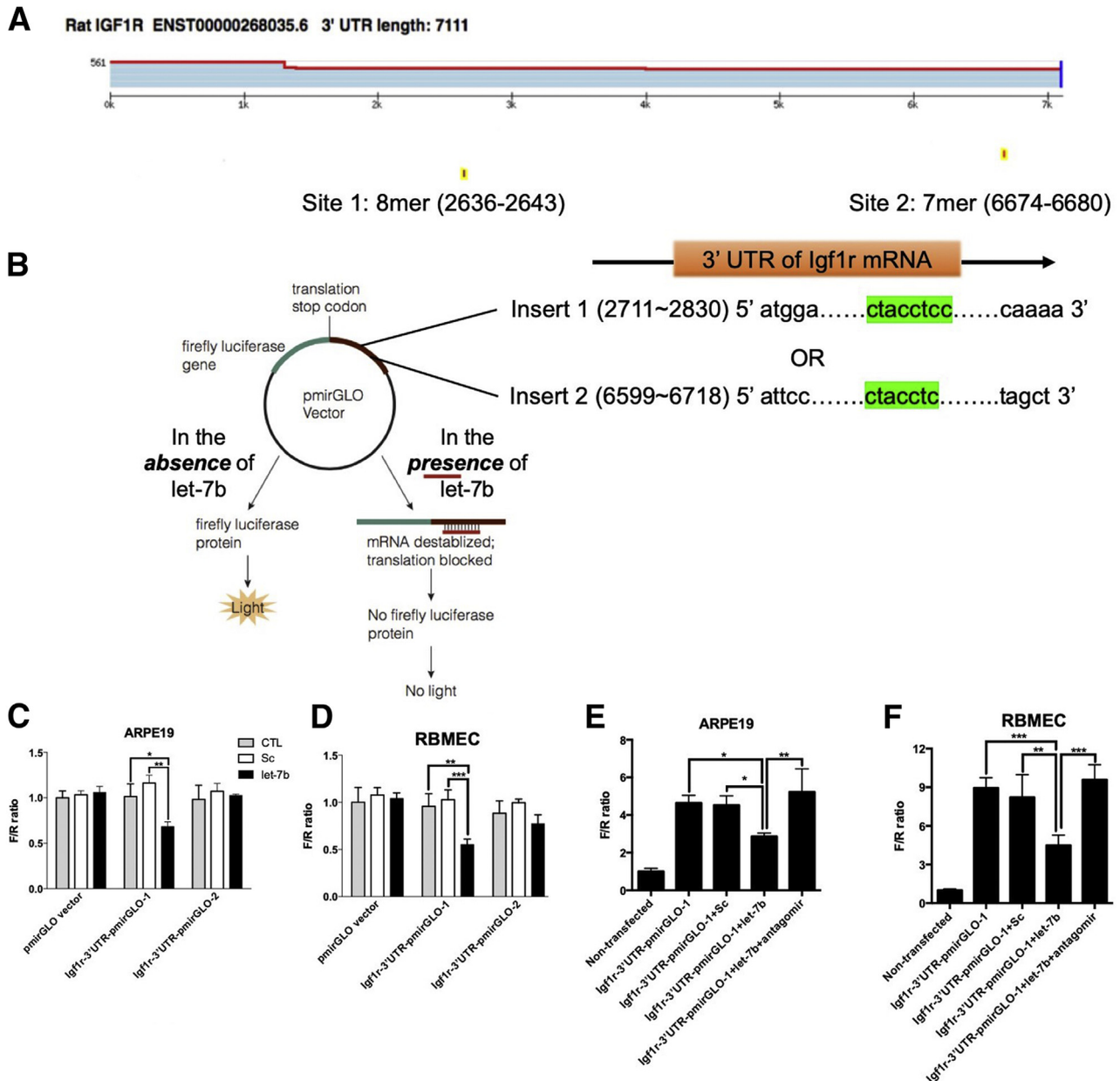


Figure 6 Dual-luciferase reporter assay confirms a let-7b target site at the 3'-untranslated region (3'UTR) of insulin-like growth factor 1 receptor (Igf1r) mRNA. **A:** *In silico* prediction of miRNA targets, based on TargetScan version 6.2, reveals two potential let-7b target sites at the 3'UTR of *Igf1r* mRNA: an 8-mer at 2636 to 2643 (site 1) and a 7-mer at 6674 to 6680 (site 2) (yellow highlights). **B:** The constitution of the reporter plasmid: a fragment of Igf1r 3'UTR that contains one of the two predicted target sites was cloned downstream of the firefly luciferase gene. Both the firefly luciferase gene and the 3'UTR fragment are transcribed into mRNA. In the presence of let-7b and a genuine let-7b target site, the mRNA is destabilized, or its translation is blocked, leading to a decrease in luciferase signal. The green highlighting shows the inserted sequences (1 and 2). **C** and **D:** In both ARPE19 and rat brain microvascular endothelial cells (RBMECs), cotransfection with let-7b mimics (at 20 nmol/L) abrogated luciferase signals in those with plasmids containing target site 1 (Igf1r-3'UTR-pmirGLO-1) but not site 2 (Igf1r-3'UTR-pmirGLO-2). **E** and **F:** In the presence of anti-let-7b antagomir, the firefly luciferase signal was restored in cells that received Igf1r-3'UTR-pmirGLO-1 and let-7b. Values in histograms are of firefly OD/Renilla OD (F/R) ratio (see *Materials and Methods*). Data are expressed as means \pm SEM (**C–F**). $n = 3$ separate experiments (**C–F**). * $P < 0.05$, ** $P < 0.01$, and *** $P < 0.001$ for indicated comparisons. CTL, control; SC, scrambled.

effective in decreasing *p53* expression and rescued *Igflr* expression (Supplemental Figure S5, B and C). More important, siRNA969 (5 ng/mL) injected subretinally (at P14) into eyes of animals subjected to OIR efficiently decreased *p53* and *Cdkn1a* mRNA levels at P18 (Figure 4, A–D) and rescued *Igflr*, *Igf1*, and *Vegfa*

expression in choroid and RPE (Figure 4, E–L); accordingly, siRNA969 treatment rescued central choroidal thickness in OIR subjects (Figure 4, M and N), along with an increase in proliferating cell nuclear antigen reactivity, indicative of cell proliferation in choroid (Supplemental Figure 5D).

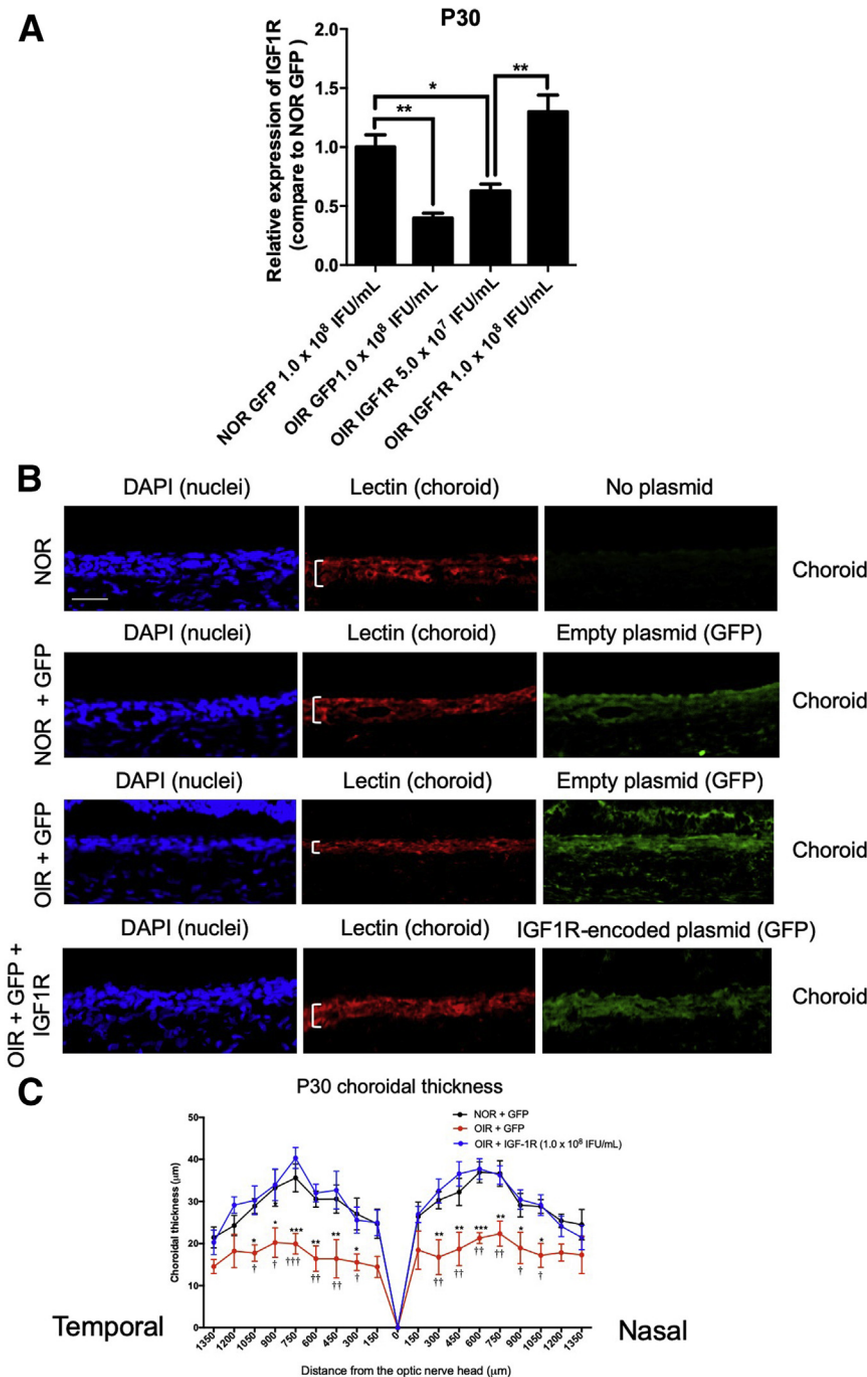


Figure 7 Subretinal restoration of insulin-like growth factor 1 receptor (*Igf1r*) expression rescues choroidal thickness. **A:** A pilot experiment was performed to identify effective dose of IGF1R-encoded lentivirus to renormalize *Igf1r* expression in the subretina; lentivirus encoding green fluorescent protein (GFP) was used as a control. Animals were injected subretinally at postnatal day (P) 14 and tissues were collected at P30. Lentivirus encoding IGF1R at 10^8 infection units (IFU)/mL (but not a lower dose) effectively normalized *Igf1r* expression in retinal pigment epithelium/choroid complex, comparable to that observed in normoxia (NOR). Values are of *Igf1r* fold change in mRNA relative to NOR. **B:** Representative images of central choroid (approximately 450 μm away from the optic nerve) are shown. Blue, red, and green signals represent nuclei (DAPI), choroid (lectin), and GFP (or IGF1R-GFP), respectively. The brackets outline choroidal thickness. **C:** Choroid thickness (the spider graph) assessed from the optic nerve to nasal and temporal periphery in groups indicated. Data are expressed as means \pm SEM (**A** and **C**). $n = 5$ separate experiments (**A** and **C**). * $P < 0.05$, ** $P < 0.01$, and *** $P < 0.001$ compared with corresponding NOR or as for indicated comparisons; $^{\dagger}P < 0.05$, $^{\dagger\dagger}P < 0.01$, and $^{\dagger\dagger\dagger}P < 0.001$ compared with corresponding oxygen-induced retinopathy (OIR) animals injected with IGF1R-encoded lentivirus. Scale bar = 50 μm (**B**).

p53 Down-Regulates *Igf1r* Expression by Inducing Let-7b miRNA

To determine how p53 governs the expression of *Igf1r* in OIR and given that numerous factors are concomitantly affected (Figure 1E), the effect of OIR in altering expression of miRNAs and the role of p53 in governing these miRNAs were evaluated.^{19,22,44} Using an 84 miRNA PCR array to profile miRNAs involved in vasoproliferation, eight miRNAs (namely, miR-

107-3p, miR-192-5p, miR-429, let-7b-5p, let-7d-5p, miR-200a-3p, miR-203a-3p, and miR-29b-3p) exhibited statistically significant changes by OIR (Figure 5A and Table 4). Of interest, only let-7b-5p (referred as let-7b hereafter), recently implicated in the inhibition of choroidal neovascularization in diabetic retinopathy,⁴⁵ was consistently found to be elevated in a similar way as p53 (from P3 to P18) in choroid and RPE of OIR-exposed rats (Figure 5, B and C), suggesting a potential regulatory interaction between p53 and let-7b; the other

miRNAs showed an irregular and an inconsistent pattern relative to p53 in RPE or in choroid throughout the period analyzed from P3 to P18 (compared with [Supplemental Figure S6](#)). Moreover, miRNA profiling in OIR in rats using next-generation sequencing revealed let-7b as one of the most expressed miRNAs in choroid.⁴⁶ In line with these observations, expression of the p53-regulated miRNA binding protein *Lin28*,¹⁵ which suppresses let-7b function,⁴⁷ was found to be decreased in OIR ([Supplemental Figure S7](#)). Although other potential p53 downstream pathways were considered,¹⁹ these did not seem to be relevant herein, as was the case for VEGF receptor 1 and hypoxia-inducible factor-1 α , which were unaltered by OIR and/or PFT α ([Supplemental Figure S7](#)). Thus, because let-7b is an important regulator of Igf1r in cancer cells,^{48,49} itself controlled by p53 ([Figure 4](#)), the link between p53 and let-7b was further corroborated by p53 knock-down (using siRNA969), which normalized *let-7b* expression in choroid and RPE of animals subjected to OIR at P14 ([Figure 5](#), D and E).

Confirming the Igf1r Transcript Is a True Target of Let-7b

On the basis of evidence that let-7 family miRNAs target the Igf1 pathway in cancer and diabetes,^{23,50} this evidence was confirmed in RPE and neurovascular endothelial cells. *In silico* prediction of potential miRNA targets using TargetScan version 6.2 revealed two potential let-7b target sites on the 3'UTR of *Igf1r* mRNA: an 8-mer at 2636 to 2643 (site 1) and a 7-mer at 6674 to 6680 (site 2) ([Figure 6A](#)). These sequences were inserted downstream of luciferase firefly gene into a pmirGLO vector ([Figure 6B](#)); individual constructs and synthetic let-7b mimic (Syn-rno/hsa-let-7b-5p, effective at 20 nmol/L) ([Supplemental Figure S8](#)) were then transfected into ARPE-19 and neurovascular endothelial cells. In the presence of let-7b mimics, luciferase activity (measured as the firefly OD/Renilla OD ratio) diminished in the groups receiving the construct that contains target site 1 (2636 to 2643) but not site 2 (6674 to 6680), indicating the former as the valid target site of let-7b ([Figure 6](#), C and D); the scrambled sequence did not alter *Igf1r* expression. In addition, cotransfection of let-7b mimic with its antagomir preserved luciferase activity ([Figure 6](#), E and F). These data collectively indicate that the *Igf1r* mRNA is targeted by let-7b.

Restoration of Igf1r Expression Rescues Choroidal Thickness after OIR

As choroidal thickness remains sustained beyond the high oxygen exposure, it was determined if restoring the levels of Igf1r in the subretina could rescue choroidal thickness. Because the amount of let-7b antagomirs needed for *in vivo* experiments was calculated to be high and possibly result in undesired effects, Igf1r was subretinally restored in OIR and it was determined if the latter would reinstate normal choroidal thickness. Lentiviral particles encoded with green

fluorescent protein (negative control) and/or Igf1r were injected into the subretinal space at low [5.0×10^7 infection units (IFU)/mL] or high (1.0×10^8 IFU/mL) doses in OIR rats at P14. Two weeks later (at P30), the expression level of Igf1r was evaluated by real-time quantitative PCR and immunohistochemistry ([Figure 7A](#) and [Supplemental Figure S9](#)). Choroidal thickness was also assessed. The higher dose of lentivirus encoded with Igf1r increased receptor expression in the subretina ([Figure 7B](#)) and rescued choroid thickness ([Figure 7](#), B and C). Hence, Igf1r exerts a significant role in reinstating choroidal vascular integrity. Altogether, inhibiting p53/let-7b pathway and/or restoring the levels of Igf1r in the subretina can avert choroidal involution and reestablish its integrity ([Supplemental Figure S10](#)).

Discussion

The inner retina has long been considered to be the primary region affected in human ROP and animal models.² Lately, dysfunctions of the outer retina and subretina, composed of photoreceptors, RPE, and choroid, have been detected in a significant number of older children formerly afflicted with ROP.^{8,51,52} Choroidal involution was confirmed in the OIR model⁶ and resulted in hypoxic ischemia, which, in turn, compromised photoreceptor integrity.⁷ Prompt revascularization is considered a desirable physiological response to ischemia to prevent long-term damage.⁵³ Strikingly and unlike the retinal (and brain) vasculature,² choroidal revascularization is negligible and involution is sustained long after OIR/ROP affliction.^{3–6} IL-1 β contributes significantly to choroidal thinning, and early anti-IL-1 β treatment (during OIR induction) prevents choroidal involution⁷; however, late anti-IL-1 β intervention (starting at P60) fails to rescue an involuted choroid.⁷ In the present study, for the first time, molecular mechanisms underlying the curtailed ability of the choroid to revascularize after OIR/ROP-induced choroidal thinning were reported. These findings reveal evidence for oxidative and possibly hypoxic-ischemic stress-related replicative senescence in OIR associated with up-regulation of cell fate-relevant p53 and downstream *let-7b* miRNA in the subretina, resulting in suppression of major permissive growth factor Igf1r (and its ligand Igf1); p53 inhibition or gene silencing enhanced choroidal vasoproliferation as it depressed let-7b expression, normalized Igf1r, and restored choroidal thickness, as observed with *Igf1r* gene rescue.

A salient feature of this study applies to the role of p53 in ROP/OIR, which has never been addressed before, particularly in the context of the subretina. Herein, we imply involvement of p53 in sustained choroidal thinning and inability to recover choroidal integrity. It was first shown that an important trigger of p53 expression (specifically, oxidative stress) was detected in the subretina, as witnessed by early up-regulation of NADPH oxidase (superoxide-

producing enzyme) transcripts, suppression of SOD, and an increase in oxidative stress—response transcription factor Nfe2l2 [alias nuclear respiratory factor 2 (Nrf2)].⁵⁴ Moreover, NOXs and Nrf2 remained increased 3 to 6 days after supplemental oxygen exposure. Oxidative stress stabilizes and activates p53,^{10,55} along with Cdkn1a.⁵⁶ Yet, one cannot exclude a role for choroid/RPE hypoxic ischemia in p53-related senescence as a result of choroidal thinning.⁷ Second, pharmacologic inhibition (PFT α) and gene knock-down of p53 restored choroidal endothelium's ability to sprout and increased choroid thickness and revascularization (indicated by increased proliferating cell nuclear antigen). These findings are consistent with effects of Nutlin-3, which is p53 pathway dependent, in inhibiting hind limb and intravitreal retinal neovascularization.⁵⁷ Third, oxidative stress preceded increased p53 expression as the two are tightly linked to cellular senescence¹⁶; hypoxic ischemia may also participate in this cell senescence after P6.

The p53 lies unequivocally at the crossroads of cell fate, primarily homeostatic repair and senescence.⁵⁸ Mechanisms for distinct roles of p53 on diverse cell fates appear to imply differences in interactions of the various isoforms of p53 with distinct target proteins involved in cell fate⁵⁹ and coinduction of epigenetic regulatory mechanisms in response to original stressors.⁶⁰ In line with its role in senescence, p53 can interfere with actions of various growth-promoting factors,⁶¹ including induction of the Igf1r pathway.⁶² It is also suggested that transient activation of p53 in response to a low dosage of oxidative stress favors senescence, whereas persistent activation of p53 during massive oxidative stress can result in cell death⁶³; yet notably, apoptosis was not detected in the choroid of OIR-subjected animals.

Of interest, the choroid and RPE exhibited similar patterns of *p53*, *Cdkn1a*, *let-7b*, *Igf1r*, and *Igf1* expression in animals subjected to OIR. Moreover, these factors also seem to be similarly regulated by p53 in choroid and RPE. In this context, both replicative and post-mitotic cells can be targeted by oxidative stress and hypoxia-ischemia, as seen in OIR, resulting in an endoplasmic reticulum stress that can be a gateway to a senescence phenotype.^{64,65} Hence, given that the RPE contributes to choroidal integrity, the impact on the choroid appears dual, such that replicative senescence would directly curtail choroid thickness, whereas non-replicative senescence in RPE could additionally indirectly affect choroid function through paracrine effects.^{12,13} The senescent phenotype in RPE may thus amplify the inability of the choroidal endothelium to proliferate, compared with the revascularization that takes place in the retinal vasculature in OIR/ROP. In this context, a role for anti-inflammatory cytokines, such as IL-10 and substance P, may also participate.^{66,67}

Recent efforts have been made to identify crucial miRNA-target pairs that regulate retinal neovascularization.^{68,69} Among them, the let-7 family of miRNAs has been detected in the plasma of patients with wet

age-related macular degeneration⁷⁰ and is down-regulated in retinal endothelium of streptozotocin-induced diabetic rats.⁷¹ miRNA profiling in choroid of OIR-subjected rats highlights the abundance of let-7b,⁴⁶ consistent with current observations. In addition, the luciferase reporter system confirms the let-7b target site on the 3'UTR of *Igf1r* mRNA (2636 to 2643) in cultured RPE and neurovascular endothelial cells.

Although the interaction between let-7 and IGF1R has been described in cancer and diabetes,^{23,50} the p53-let-7b-Igf1r regulatory axis is new to retinopathies. It has previously been shown that p53 does not directly bind to *Igf1r* promoter,⁷² but rather regulates the Igf1 pathway by modulating antiproliferative miRNAs.¹⁹ The let-7 family stands out particularly intertwined with the Igf1 axis.^{23,48,73} Biogenesis of the let-7 family is enhanced by p53, which represses Lin28, an RNA binding protein that precludes let-7 maturation and action.^{22,74,75} Consistent with increased p53 and let-7b in OIR, Lin28 was suppressed.

At first glance, down-regulation of prosurvival/angiogenic Igf1r in the subretina in OIR as the choroid thins is paradoxical, especially in ischemic retinopathy, where retinal neovascularization is driven by ischemia-hypoxia. Igf1r, which is ubiquitously expressed, is linked to elevated metabolism,⁷⁶ contributes to oxidative stress generation, and affects longevity.^{77,78} Thus, a reduction in Igf1r, as observed herein, would reduce metabolic rate in OIR,⁷⁹ and intuitively be considered beneficial. On the other hand, on IGF1 stimulation during hypoxia,^{80,81} RPE releases proangiogenic/prosurvival IGF1/2, IGF-binding proteins 3/6 (key binding proteins for bioactive IGF1/2), and VEGFa. Conversely, low postnatal serum Igf1 values have been strongly correlated with development of ROP⁸²; but supplementing preterm infants with IGF-1 failed to prevent ROP (clinical trial: NCT01096784), possibly because of limited Igf1r expression in the inner retina. Low Igf1r, Igf1, and Vegfa expression is also observed in the subretina of OIR subjects; a defect in RPE-derived VEGF can lead to choriocapillaris atrophy and deleterious choroidal remodeling,⁸³ which share pathologic features with middle-aged OIR/ROP subjects (notably, choroidal degeneration, abnormalities in RPE, and photoreceptor dysfunction).⁷ All in all, a reduction in the expression of Igf1, Igf1r, and Vegfa, as seen in the outer retina of OIR subjects, generates an unfavorable environment for choroidal revascularization, resulting in persistent choroidal involution; decreased Igf1r signaling in the RPE may further compromise photoreceptor function.^{7,8,84}

Collectively, this study reveals an unprecedented mechanism to explain the inability of the choroid to revascularize after OIR-induced involution. This study unveils, for the first time, the critical role of p53-let-7b-Igf1r axis in orchestrating a defective angiogenic environment in the subretina, which impedes choroidal revascularization. Targeting this pathway represents a potential therapeutic avenue to rescue choroidal involution in ROP subjects and other chorioretinopathies, as in geographic atrophy.

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Supplemental Data

Supplemental material for this article can be found at <http://doi.org/10.1016/j.ajpath.2019.07.009>.

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