Enhancing Retinal Endothelial Glycolysis by Inhibiting UCP2 Promotes Physiologic Retinal Vascular Development in a Model of Retinopathy of Prematurity

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PURPOSE. We address the hypothesis that uncoupling protein 2 (UCP2), a cellular glucose regulator, delays physiologic retinal vascular development (PRVD) by interfering with glucose uptake through glucose transporter 1 (Glut1).

METHODS. In the rat 50/10 oxygen-induced retinopathy (OIR) model, retinal Glut1 and UCP2 were measured and compared to room air (RA)-raised pups at postnatal day 14 (p14). Pups in OIR and RA received intraperitoneal genipin, an UCP2 inhibitor, or control every other day from p3 until p13. Analyses at p14 included avascular/total retinal area (AVA), Western blots of retinal UCP2 and Glut1, and immunostaining of Glut1 in retinal cryosections. Intravitreal neovascular/total retinal area (IVNV) was analyzed at p18, and electroretinograms were performed at p26. Glut1 and phosphorylated VEGFR2 (p-VEGFR2), glucose uptake, adenosine triphosphate (ATP) production, and cell proliferation were measured in human retinal microvascular endothelial cells (hRMVECs) pretreated with genipin or transfected with UCP2siRNA, Glut1siRNA, or control siRNA when incubated with VEGF or PBS.

RESULTS. At p14, OIR pups had increased AVA with decreased Glut1 and increased UCP2 in the retina compared to RA retinas. Intraperitoneal genipin increased retinal Glut1 and reduced AVA. Compared to control, treatment with genipin or knockdown of UCP2 significantly increased Glut1, glucose uptake, ATP production, VEGF-induced p-VEGFR2 and cell proliferation in hRMVECs. Knockdown of Glut1 inhibited VEGF-induced p-VEGFR2. Genipin-treated OIR pups with decreased AVA at p14 had reduced IVNV at p18 and increased amplitudes in a- and b-waves at p26.

CONCLUSIONS. Extending PRVD by increasing retinal endothelial glucose uptake may represent a strategy to prevent severe retinopathy of prematurity and vision loss.

Keywords: angiogenesis, glucose transporter 1, oxygen induced retinopathy, VEGFR2, retinopathy of prematurity
MATERIALS AND METHODS

Animals and Ethical Statement

Timed-pregnant Sprague-Dawley rats were purchased from Charles River (Wilmington, MA, USA). Animals had free access to food and water ad libitum and were kept on a 12-hour/12-hour light-dark cycle. All animal experimental protocols were approved by the Institutional Animal Care and Use Committee of University of Utah. All experiments were carried out in accordance with the Guidelines and Regulations for the Care and Use of Laboratory Animals by the University of Utah (Salt Lake City, UT, USA) and the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Vision Research.

Rat Oxygen-induced Retinopathy (OIR) Model and Intraperitoneal Injections

The rat OIR model used in this study has been described previously.18,19 Within 6 hours of birth, newborn Sprague-Dawley pups (male and female) and dam were placed into an oxygen chamber (OxyCycler, Biospherix, NY, USA) with controlled oxygen concentrations that cycled between 50% and 10% every 24 hours. After 14 days, pups were moved into room air (RA). Each litter had 12 to 14 pups to maintain consistency in measured outcomes in the model. Experiments were repeated in four individual litters.

In a litter, pups were randomly assigned to two groups and treated with intraperitoneal injections of genipin (10 mg/kg, Sigma-Aldrich Corp., St. Louis, MO, USA) dissolved in dimethyl sulfoxide (DMSO) and PBS (1:500) or an equal volume of vehicle (DMSO+PBS) as control beginning at postnatal day 3 (p3) and repeated every other day until p13. For pups in the OIR model, injections were performed at the end of 50% oxygen cycles and before 10% oxygen cycles to avoid interventions during the 10% oxygen cycle.

Ganzfeld Electroretinogram (ERG)

Retinal function was assessed by Ganzfeld ERG (LKC Technologies, Inc., Gaithersburg, MD, USA). Pups at p20 were fully dark-adapted overnight and subsequently handled under dim red light illumination. After anesthesia and pupil dilation with tropicamide (1%) eye drops (Bausch and Lomb, Tampa, FL, USA), pups were kept on a heating pad throughout the procedure to maintain body temperature. Subdermal needle electrodes were inserted underneath the skin at the base of the tail (ground electrode) and between the eyes on the forehead (reference electrode). Corneas were lubricated with coupling gel (GenTeal, hyromellose 0.3%; Alcon, Fort Worth, TX, USA); the objective electrode was advanced near the corneal surface, and deep red illumination was used to focus on the retina. Responses to light stimuli were elicited at luminances ranging from −4.5 to 2.2 cd s m⁻² with 30 sweeps at 2-second intervals. Data were displayed as the mean amplitude (in μV) of the a-wave (as a measure of photoreceptor function) and b-wave (as a measure of bipolar cell function).

Retinal Flat Mount Preparation, Imaging, and Analysis of Peripheral Avascular Retinal Area/Total Retinal Area (AVA) and Intravitreal Neovascularization/Total Retinal Area (IVNV)

After enucleation, eyes were fixed in 4% paraformaldehyde (PFA; EMS, Hatfield, PA, USA) for 2 hours and placed into PBS (containing in mM NaCl 137, KCl 2.7, Na₂HPO₄ 8, KH₂PO₄, pH 7.4). To dissect the retina in each eye, the cornea, lens, and vitreous were removed and the RPE, choroid, and sclera were removed from the eyecup. Retinas were flattened onto slides. The retinal flat mounts were stained with Alexa-Fluor 568-labeled isoleucin GS-IB4 from Griffonia simplicifolia (2.5 μg/mL; Invitrogen, Grand Island, NY, USA) to stain the retinal vasculature. Flat mounts were imaged at ×4 magnification using an inverted fluorescence microscope (Olympus IX81; Olympus Corp, Tokyo, Japan), and individual images were joined using Metamorph 7.0 software (Molecular Devices, Inc., Sunnyvale, CA, USA). The percent of AVA and of IVNV was determined by two masked reviewers using Image J software (National Institutes of Health [NIH], Bethesda, MD, USA).

Retinal Cryosection Preparation and Staining

Anterior segments were removed from eyes after 30 minutes of fixation in 4% PFA. Eye cups were replaced into 4% PFA for an additional 1 hour and incubated overnight in 30% sucrose. After immersion in optimal cutting temperature compound (OCT; Tissue Tek; EMS, Hatfield, PA, USA), retinas were cut into 12 μm cryosections. After rinsing and blocking in blocking buffer with 5% goat serum for 1 hour at room temperature, sections were incubated with mouse anti-Glut1 (1:200; Abcam, Cambridge, MA, USA) overnight at 4°C, followed by incubation for 1 hour with FITC-conjugated goat anti-mouse (1:200; Jackson ImmunoResearch Laboratories, Inc., West Grove, PA, USA), isoleucin B4 (lectin, 1:500; Invitrogen) and TO-PRO-3 iodide (1:500; Invitrogen) to stain nuclei. Sections stained without primary antibodies were used as controls. Images were captured using confocal microscopy (Olympus IX81, Tokyo, Japan) at ×20 magnification.

Cell Culture and Transfection

Human retinal microvascular endothelial cells (hRMVECs; Cell Systems, Kirkland, WA, USA) were maintained in endothelial growth medium (EGM-MV; Lonza, Walkersville, MD, USA) supplemented with 5% FBS. Cells of passages 3 to 5 were used for experiments. To knock down glucose transporter 1 (Glut1) or UCP2, hRMVECs at 70% confluence were transfected with siRNAs targeting human Glut1 gene (solute carrier family 2 member 1, SLCC2A1) or human UCP2 (Applied Biosystems, Foster City, CA, USA) using lipofectamine 2000 or 3000 (Invitrogen). A silencer selective negative control siRNA was used as control for Glut1 or UCP2 knockdown. At 24 to 36 hours after transfection, cells were treated and used for different assays.

Glucose Uptake Assay

Human RMVECs were plated into 96-well plates at 1 × 10⁴ cells/well and grown for 24 hours. Cells then were grown in serum-free endothelial basal medium (EBM) overnight. After three washes in warm PBS, cells were incubated in Krebs-Ringer-Phosphate-HEPES (KRPH) buffer (20 mM HEPES, 5 mM KH₂PO₄, 1 mM MgSO₄, 1 mM CaCl₂, 136 mM NaCl, and 4.7 mM KCl, pH 7.4) with 2% BSA for 30 minutes. Cells then were treated with genipin (10 μM) for 2 hours before incubation with VEGF (20 ng/mL). Glucose uptake was measured after incubating with 2-NBDG (2-NBDG; 2 mM) for 30 minutes. DMSO and PBS were used as treatment controls for genipin and VEGF, respectively. After treatment, cells were washed in PBS three times to remove 2-NBDG that was not taken up, and then cells were measured for fluorescence at excitation/emission maxima of approximately 520/480 nm.
Upregulating Glut1 Promotes Retinal Vascularization

ATP Production Assay

Human RMVECs were plated into 6-well plates at 0.5 × 10⁶ cells/well and grown for 48 hours until confluence. Cells were then starved in EBM medium without serum and growth factor for 3 hours and then treated with genipin (10 μM) for 2 hours. ATP concentration was measured by fluorometric ATP assay kit (Abcam) in 96-well plates.

Cell Proliferation Assay

Human RMVECs were plated into 96-well plates at a density of 5000 cells/well. At 24 hours after plating, cells were starved in serum-free media overnight, and then pretreated with genipin (10 μM) for 2 hours before incubation with VEGFA (20 ng/mL) or control PBS for another 24 hours. Cell number was measured with the Vybrant MTT cell proliferation assay kit (Invitrogen).

Western Blots

Cells or retinas were lysed in RIPA buffer with protease inhibitor cocktail (Sigma Aldrich Corp.) and orthovanadate (Thermo Fisher Scientific, Grand Island, NY, USA). Lysates were clarified by centrifugation at 16.1 × 1000g for 5 minutes at 4°C. Protein concentration in the supernatant was quantified by the bicinchoninic acid assay (BCA). A total of 20 μg of protein from each treatment was loaded into NuPAGE 4% to 12% Bis-Tris Gels (Invitrogen), transferred to a polyvinylidine fluoride (PVDF) membrane, and incubated with antibodies to phospho-VEGFRI (1:1000, Abcam) at 4°C overnight, transferred to a polyvinylidine fluoride (PVDF) membrane, and incubated with antibodies to phospho-VEGFRI (1:1000, Abcam) at 4°C overnight. After incubating with primary antibodies, membranes were incubated with horseradish peroxidase conjugated goat anti-rabbit secondary antibody or goat anti-mouse secondary antibody (1:3000–5000, ThermoFisher Scientific) at room temperature for 1 hour. All membranes were reprobed with horseradish peroxidase conjugated α-actin (1:3000, Santa Cruz Biotechnology) as loading controls. Quantification of densitometry was performed using the licensed software UN-SCAN-IT version 7 (Silk Scientific, Inc., Orem, UT, USA).

Statistical Analysis

A mixed effects linear regression model was used to statistically analyze data collected from OIR rats for AVA and IVNV using STATA-14 software (StataCorp LLC, College Station, TX, USA). That model included fixed and random effects to account for biologic variation between litters by comparing experimental to control groups inside the same litter. The intraperitoneal injections used in this study did not contribute much variation between eyes. Only one eye from each animal was used for AVA and IVNV analysis and the fellow eye was used for protein and RNA analysis. A 2-tailed t-test was used for protein analysis and in vitro studies. Results were displayed as mean ± SEM. A minimum value of P < 0.05 was considered statistically significant. For animal studies, 16 individual pups from at least four litters were analyzed for AVA or IVNV. For ERG analyses, four to six pups were used. For immunolabeling in retinal cryosections, three to four retinal sections at 60-μm intervals were taken from one eye of three pups in each group. For protein analyses, each treatment included at least five different animals from three litters. For in vitro studies, each experimental condition included an n = 4-9 from at least two independent experiments.

RESULTS

Delayed PRVD Associated With Decreased Retinal Glut1 in OIR

As a model of human ROP, pups in the rat OIR model have incomplete PRVD at p14 in contrast to RA raised pups that have fully vascularized retinas (Fig. 1A). Endothelial cells use and produce energy locally through glycolysis to regulate endothelial cell proliferation, migration, and tip cell formation, which are cellular events involved in angiogenesis. The retina takes up glucose mainly through Glut1. Studies have shown that Glut1-mediated glucose transport is essential for postnatal development of the murine brain vasculature. Therefore, we postulated that delayed PRVD in OIR pups was associated with insufficient glucose uptake. To study this, Glut1 was first measured in retinal lysates from RA and OIR pups at p14. As shown in Figures 1B and 1C, compared to RA, Glut1 protein was significantly decreased in OIR. To determine if Glut1 is localized to retinal vascular EC, immunohistochemistry (IHC) for Glut1 was performed in retinas of p14 RA or OIR pups. IHC of Glut1 in retinal cryosections was mainly located in the retinal ganglion cell layer, inner plexiform layer, photoreceptor outer segments, and RPE. Glut1 also strongly colocalized with isolectin-stained retinal vascular ECs (Fig. 1D). Taken together, these data supported the notion that reduced Glut1 in retinal vascular ECs was associated with delayed PRVD in p14 OIR pups.

Decreased Retinal Glut1 is Associated With Increased UCP2 in OIR Pups

Mitochondrial UCP2 has been implicated in cellular glucose metabolism related to pathologic conditions, such as diabetic retinopathy. Knockdown of UCP2 significantly increases ATP production from glucose metabolism in human hepatocarcinoma cells. To determine if decreased retinal Glut1 is related to UCP2 expression, UCP2 protein was measured in the retinas by Western blots. Compared to RA, UCP2 was significantly increased in retinal lysates of OIR pups (Fig. 2) that had decreased Glut1 (Fig. 1B), supporting the thinking that lower Glut1 in OIR pups could be explained by increased UCP2.

Inhibition of UCP2 Extends PRVD by Upregulating Retinal Glut1

Since reduced Glut1 was associated with delayed PRVD and increased UCP2, we predicted that OIR increased retinal UCP2, which then decreased Glut1 and delayed PRVD. To test this, OIR pups were given intraperitoneal injections of genipin, an UCP2 inhibitor, or solvent control every other day from p3 to p15 during the 50% oxygen cycles. As shown in Figure 3, compared to control, inhibition of UCP2 by genipin (Figs. 3A, 3B) significantly increased Glut1 protein in the retinas of OIR pups (Figs. 3C, 3D). We then studied if this increased Glut1 had a role in extending PRVD. We predicted that genipin treatment would extend PRVD, reflected by lower AVA. Compared to control, genipin-treated pups had significantly decreased AVA measured in isolectin-stained flat mounts at p14 (Figs. 3E, 3F), supporting the prediction.

To further test the effects of inhibition of UCP2 in extending PRVD by upregulating Glut1, a series of experiments were performed in hRMVECs. We first asked if genipin treatment would increase Glut1 and, therefore, promote glucose uptake.
to increase ATP production. To test this hypothesis, we measured Glut1 protein by Western blot, glucose uptake by (N-[7-Nitrobenz-2-oxa-1,3-diazol-4-yl] Amino)-2-Deoxyglucose and ATP production by the fluorometric assay. Treatment with the UCP2 inhibitor, genipin, for 2 hours significantly increased Glut1 protein (Figs. 4A, 4B). To eliminate potential off-target effects of genipin treatment, Glut1 protein was measured in hRMVECs transfected with UCP2siRNA. As shown in Figures 4C and 4D, knockdown of UCP2 by siRNA significantly increased Glut1 protein, similar to genipin treatment. Genipin treatment for 30 minutes significantly increased glucose uptake (Fig. 4E) and ATP production (Fig. 4F), supporting the thinking that inhibition of UCP2 increases Glut1 protein and its mediated glucose uptake.

Variable oxygen reduced retinal VEGFR2 in rats in the OIR model compared to those in RA. Activation of extracellular signal-responsive kinase-1/2 (ERK1/2), a downstream signaling effector of VEGF, was significantly reduced in the retina of rat pups in OIR compared to those raised in RA (Supplemental Figure), supporting the thinking that reduced activation of VEGF signaling contributes to decreased angiogenesis. We then determined if increased Glut1-mediated glucose uptake from genipin treatment would increase the angiogenic activity of hRMVECs. Compared to respective controls, VEGF-induced phosphorylation of VEGFR2 (p-VEGFR2) was significantly increased in genipin-treated hRMVECs (Figs. 5A, 5B) and in
hRMVECs knocked down for UCP2 by siRNA transfection compared to control (Figs. 5C, 5D). Genipin treatment increased p-VEGFR2 and hRMVEC proliferation (Fig. 5E). To determine if increased p-VEGFR2 from genipin treatment was dependent on Glut1, p-VEGFR2 was measured in hRMVECs transfected with Glut1 siRNA or control siRNA. Compared to control siRNA-transfected hRMVECs, knockdown of Glut1 by Glut1 siRNA reduced VEGF-induced p-VEGFR2 in hRMVECs (Fig. 5F). Taken together, the data demonstrated that inhibition of UCP2 by genipin increases Glut1 and its mediated glucose uptake and, therefore, promotes angiogenic activity of hRMVECs through activation of VEGFR2.

Genipin-Induced PRVD Reduces IVNV and Improves ERG

Extending PRVD reduces hypoxic retina available to stimulate pathologic angiogenesis. We predicted that genipin-treated OIR pups with less AVA, an indicator of extended PRVD, would have less IVNV. Compared to control-treated OIR pups, genipin treatment from p3 to p13 significantly reduced IVNV (Figs. 6A, 6B). To address the question if the reduced IVNV at p14 was related to reduced AVA at p14 rather than genipin treatment, IVNV was measured in OIR pups treated with genipin or control every other day from p14 to p18, instead of from p3 to p13. As shown in Figures 6C and 6D, compared to control/control treatment, late genipin treatment did not reduce IVNV in the OIR pups, supporting the thinking that increased PRVD at p14 from early genipin treatment accounted for the subsequent reduced IVNV.

Since genipin treatment extended PRVD, we predicted that it also would improve retinal function measured by ERG. At p26, OIR pups have lower a- (Fig. 7A) and b-wave (Fig. 7B) amplitudes compared to RA, consistent with other studies.24,25 Compared to respective controls, genipin treatment significantly increased a- (Fig. 7A) and b-wave (Fig. 7B) amplitudes in OIR but not in RA pups. The OIR pups treated with genipin still maintained less AVA (Fig. 7C) and IVNV (Fig. 7D) compared to controls; however, the difference was not significant probably because the rat OIR model undergoes natural regression of IVNV and extension of PRVD.

Taken together, the data provided evidence that pups treated with genipin experience extended PRVD, reduced IVNV, and improved neural retinal function.

DISCUSSION

The focus of treatment in ROP has been to inhibit pathologic intravitreal neovascularization. However, recurrent intravitreal neovascularization after anti-VEGF treatment occurs in clinic,
and animal studies. Therefore, only inhibiting intravitreal neovascularization without extending PRVD may not be an optimal treatment for ROP, because persistent avascular retina stimulates hypoxia-induced recurrent intravitreal neovascularization and impairs vision function due to loss of retinal vasculature. We studied the role of glucose uptake and energy metabolism in retinal vascular endothelial cells needed to extend PRVD and addressed the hypothesis that UCP2, a cellular glucose regulator, extended PRVD by interfering with glucose uptake through Glut1.

The retina is a metabolically active organ with high-energy demands. Glucose is the primary energy supply for the retina, although lipids also can be metabolized by retinal photoreceptors. The retina takes up glucose through Glut1 and generates ATP by aerobic glycolysis. Several studies have established the importance of glucose metabolism in retinal neuronal survival and neurovascular interactions. Recently, there is interest in the role of glycolysis in angiogenesis particularly in diseases involving aberrant vessel growth. Pharmacologic blockade of the glycolytic activator, phosphofructokinase-2/fructose-2,6-bisphosphatase 3 (PFKFB3) with a small molecule, 3-(3-pyridinyl)-1-(4-pyridinyl)-2-propan-1-one (3PO) reduced retinal vascular branching and radial extension and choroidal neovascularization in a mouse laser-induced model of choroidal neovascularization, providing evidence that glycolysis-mediated energy metabolism regulates angiogenesis.

We found that delayed PRVD in OIR pups correlated with reduced retinal Glut1 due to insufficient glucose uptake to support retinal angiogenic growth. In support of this conjecture, we also observed an association between reduced Glut1 and increased retinal UCP2. UCP2 has been involved in cellular lipid and glucose metabolism. Knockout of UCP2 promotes cell proliferation by stimulating cellular use of glycolysis-derived pyruvate in mouse embryonic fibroblasts. To test the hypothesis that insufficient glucose uptake reduced glycolytic metabolism and delayed PRVD, we treated OIR pups with the UCP2 inhibitor, genipin, and found inhibition of UCP2 increased retinal Glut1 protein in association with reduced AVA at p14. We also observed that in OIR, VEGF was increased but activation of the downstream signaling effector of VEGFR2, p-ERK, was decreased at p14. Maintaining activation of the VEGF signaling pathway requires VEGFR2 internalization by endocytosis and recycling, which are ATP-demanding cellular events. We further provided evidence that insufficient ATP production from reduced Glut1 reduced

**FIGURE 4.** Inhibition of UCP2 increases Glut1 and VEGF-induced glucose uptake in hRMVECs. Western blot of Glut1 and UCP2 (A, B) in hRMVECs treated with control or genipin (10 μM) for 2 hours and (C, D) in hRMVECs transfected with UCP2 siRNA or control siRNA. (A, C) Representative gel images. (B, D) Quantification of densitometry of Glut1. (E) Glucose uptake measured by (N-[7-Nitrobenz-2-oxa-1,3-diazol-4-yl]Amino)-2-Deoxyglucose and (F) ATP production in hRMVECs treated with genipin (10 μM) or control for 30 minutes (*P < 0.05, **P < 0.01, ***P < 0.001 versus Control; †P < 0.05 versus Control siRNA; n = 4 for Western blots and n = 9 for glucose uptake and ATP production).
VEGFR2 signaling. In hRMVECs, inhibition of UCP2 by genipin increased VEGF-induced glucose uptake, ATP production, and VEGF-mediated activation of VEGFR2, whereas knockdown of Glut1 inhibited VEGFR2 activation. Our results supported the thinking that UCP2-mediated reduction in Glut1 interferes with VEGF-induced signaling and, therefore, reduces intraretinal vascularization necessary for PRVD. The importance of Glut1 expression in retinal endothelial cells also is highlighted in a study of diabetic retinopathy, in which hyperglycemia-induced oxidative stress disrupted glucose homeostasis by reducing Glut1 plasma membrane expression. Future studies are needed to determine if oxidative stress is involved in reduced retinal Glut1 and if UCP2 quenches reactive oxygen in OIR.

We then addressed the question whether extending PRVD by genipin treatment would reduce IVNV and improve retinal function. We observed that genipin-treated OIR pups that had less AVA at p14 had decreased IVNV at p18 and increased a- and b-wave amplitudes at p26 following overnight dark adaptation. These improved ERG amplitudes were observed in OIR pups, but not in RA pups, suggesting that UCP2 activity does not regulate retinal function in normal retinal development. However, others reported that 10- to 15-week-old UCP2 knockout mice had reduced a-waves, but normal b-waves after 1 hour of dark adaptation. The differences in the studies may relate to oxygen stresses in our pups and differences in animal age or possible dark adaptation time, since overnight dark adaptation is needed to measure ganglion cell function in the ERG.43 Future studies are warranted to understand the potential role of UCP2 and glucose metabolism in promoting photoreceptor maturation, neurovascular interactions, and in extending PRVD. Interestingly, room air raised pups treated with genipin every other day from p3 to p13 showed fully vascularized retinas at p14, suggesting that UCP2 is not involved in retinal vascular development. Therefore, inhibition of UCP2 may be a potentially safe target to treat ROP.

The importance of glucose homeostasis in preterm infants is recognized in neonatal care. Some studies proposed hyperglycemia as a potential risk factor for ROP. Hyperglycemia occurs in the setting of stress induced insufficient insulin secretion or relative insulin resistance. However, the interpretation of hyperglycemia is confounded, because it occurs in very stressed premature infants who have other conditions of prematurity that may influence the risk of ROP.
Figure 6. Intraperitoneal genipin to reduce AVA reduces intravitreal neovascularization. (A) and (B) IVNV at p18 of OIR pups received intraperitoneal injections of genipin or control from p3 to p14. (A) Representative flat mount images and (B) quantification of IVNV; *$P<0.05$ versus Control. Scale bar: 990 µm. (C, D) IVNV at p18 of OIR pups treated with control or genipin from p14 to p18. (C) Representative flat mount images and (D) quantification of IVNV. Scale bar: 990 µm, $n=3–16$.

Figure 7. Intraperitoneal genipin increases retinal function. Ganzfeld ERG of (A) a-wave amplitude and (B) b-wave amplitudes in p26 RA and OIR pups after overnight dark adaptation and treatment with control or genipin from p3 to p13 (*$P<0.05$, **$P<0.001$ versus RA/Control; †$P<0.01$, ††$P<0.001$ versus OIR/Control; $n=4–6$ for ERG and $n=11$ for AVA and IVNV); and quantification of (C) AVA and (D) IVNV of OIR pups at p26 treated with control or genipin from p3 to p13.
and other clinical studies have not identified sufficient evidence to support a direct causal effect between high glucose and ROP. Preterm neonates, particularly those of young gestational ages or with intrauterine growth restriction, are prone to suffer hypoglycemia because of limited storage of glycogen and fat, inability to generate new glucose through gluconeogenesis by underdeveloped organs including the liver, and high metabolic demands from a relatively high metabolism rate. Preterm infants with hypoglycemia have abnormal neurodevelopment due to insufficient energy supply from glucose oxidation. These clinical observations emphasize the importance of glucose metabolism in preterm infants and relate to our experimental study on physiologic retinal neural and vascular development and pathologic neovascularization. In conclusion, our study supported the line of thinking that regulating uptake of cellular glucose and glycolytic metabolism of the retinal vascular endothelial cells is important to extend PRVD and reduce retinal pathology in ROP and prematurity. Furthermore, improved glucose uptake and glycolysis may improve neural function either from improved vascularity or through other mechanisms.

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