In Vivo Imaging of Retinal Hypoxia Using HYPOX-4-Dependent Fluorescence in a Mouse Model of Laser-Induced Retinal Vein Occlusion (RVO)

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Purpose. To demonstrate the utility of a novel in vivo molecular imaging probe, HYPOX-4, to detect and image retinal hypoxia in real time, in a mouse model of retinal vein occlusion (RVO).

Methods. Retinal vein occlusion was achieved in adult mice by photodynamic retinal vein thrombosis (PRVT). One or two major retinal vein(s) was/were occluded in close proximity to the optic nerve head (ONH). In vivo imaging of retinal hypoxia was performed using, HYPOX-4, an imaging probe developed by our laboratory. Pimonidazole-adduct immunostaining was performed and used as a standard ex vivo method for the detection of retinal hypoxia in this mouse RVO model. The retinal vasculature was imaged using fluorescein angiography (FA) and isolecin B4 staining. Retinal thickness was assessed by spectral-domain optical coherence tomography (SD-OCT) analysis.

Results. By application of the standard ex vivo pimonidazole-adduct immunostaining technique, retinal hypoxia was observed within 2 hours post-PRVT. The observed hypoxic retinal areas depended on whether one or two retinal vein(s) was/were occluded. Similar areas of hypoxia were imaged in vivo using HYPOX-4. Using OCT, retinal edema was observed immediately post-PRVT induction, resolving 8 days later. Nominal preretinal neovascularization was observed at 10 to 14 days post-RVO.

Conclusions. HYPOX-4 is an efficient probe capable of imaging retinal hypoxia in vivo, in RVO mice. Future studies will focus on its use in correlating retinal hypoxia to the onset and progression of ischemic vasculopathies.

Keywords: retinal hypoxia, ischemia, retinal vein occlusion, in vivo imaging, HYPOX-4
Generally, it is assumed that retinal capillary nonperfusion provides some indirect measure of retinal hypoxia, and in the clinic, this is routinely assessed by fluorescein angiography (FA). The application of this technique provides conclusive findings of retinal capillary nonperfusion in approximately 50% to 60% of RVO cases. Therefore, the distinction between ischemic and nonischemic RVO is limited, if not totally unreliable. In many cases, the confirmation, the exact location, and the duration of the venous occlusion are uncertain making it difficult to assess any associated ischemia-induced hypoxia. Oxygen-dependent molecular phosphorescence quenching, oxygen sensitive electrodes, nuclear magnetic resonance (NMR), retinal oximetry, Doppler optical coherence tomography (D-OCT), visible-light OCT (vis-OCT), and immunohistochemical analysis, have been used to measure retinal oxygen levels, but each has its own drawbacks and none are suitable methods to assess retinal hypoxia secondary to ischemia in vivo and in real time.

Molecular imaging techniques hold the potential, as powerful in vivo tools, to detect and image retinal hypoxia perhaps allowing the use of hypoxia as a biomarker for the onset, progression, and resolution of ischemic retinopathies. Assessment of retinal hypoxia has been demonstrated by application of the ex vivo pimonidazole immunostaining technique. Pimonidazole contains the nitroimidazole moiety that mimics oxygen; nitroreductase enzymes in tissues with low-oxygen pressure reduce it, presumably causing its sequestration via pimonidazole-protein adduct formation. Detection and quantification of retinal hypoxia are performed by excising the tissue of interest, and staining it with antibodies that recognize these protein adducts. Recently, we have synthesized imaging probes incorporating hypoxia sensitive moieties and fluorescent dyes, and tested their reactivity, and pharmacokinetic parameters for its application to the in vivo characterization of retinal hypoxia. To better understand the occurrence and the relationship of retinal hypoxia to the pathogenesis of RVO, we tested the ability of HYPOX-4 to image retinal hypoxia in vivo, in mouse RVO. We compared our results with those obtained by application of the standard ex vivo pimonidazole technique. Herein, we report the results.

**MATERIALS AND METHODS**

**Synthesis of HYPOX-4**

The synthesis of HYPOX-4 was discussed in our previous report.

**Animal Procedures**

C57BL/6j male mice 4 to 6 weeks of age were purchased from Charles River Laboratories (Chicago, IL, USA). All animal procedures used in this study were approved by the Vanderbilt University Institutional Animal Care and Use Committee and were performed in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

**Retinal Vein Occlusion in Mice**

A detailed method for photodynamic retinal venous thrombosis (PRVT) in mice has been described in previous publications and was replicated in this study with slight modifications. Briefly, C57BL/6 male mice were anesthetized by intraperitoneal injection of a ketamine/xylazine (25/10 mg/kg) mixture. The eyes were dilated using 1% tropicamide and Gentec lubricating eye gel (Alcon, Fort Worth, TX, USA) was used to keep the eyes moist. A PBS solution of 40 mM rose bengal (4,5,6,7-tetrachloro-2',4',5',7'-tetraiodo-fluorescein di-sodium salt, certified purity, 95%; Sigma-Aldrich Corp., St. Louis, MO, USA) was sterilized by filtration through a 0.22-μm filter. Mice received tail vein injections of rose bengal at a dose of 40 mg/kg. An argon laser photocoagulator (blue-green) mounted on a slit-lamp (Space Coast Laser, Inc., Palm Bay, FL, USA) was used to create RVO (50-μm spot size, 1-second duration, 50 mW). To completely stop the blood flow, two to three laser applications were required as determined by observing the fundus image with a slit-lamp microscope. In each fundus, one or two veins were photocoagulated to create retinal ischemia.

**In Vivo and Ex Vivo Imaging of Retinal Hypoxia Using HYPOX-4 in RVO Mice**

Two hours post-RVO, mice were injected intraperitoneally with HYPOX-4 (60 mg/kg body weight). Twenty-two hours post injection of HYPOX-4, in vivo HYPOX-4-dependent fluorescence imaging was performed. The mice were anesthetized with a ketamine/xylazine (25/10 mg/kg) mixture, dilted with 1% tropicamide, and placed on a warm platform, Gentec lubricating eye gel was used to keep the eyes moist. Fluorescent and bright field fundus images were acquired using the Micron IV retinal imaging system (Phoenix Research Laboratories, Pleasanton, CA, USA). Mice were killed, enucleated, and the globes were fixed in 10% neutral buffered formalin (NBF). Retinas were dissected, stained with Alexa Fluor 647-conjugated isoclin B4 and flat-mounted on a microscope slide with Prolong Gold mounting medium (Life Technologies, Grand Island, NY, USA). Images were captured using an epifluorescence Nikon Eclipse Ti-E inverted microscope (Nikon, Melville, NY, USA).

**Ex Vivo Detection of Retinal Hypoxia in RVO Mice Using Pimonidazole-Adduct Immunostaining**

Two hours post-RVO, mice received intraperitoneal injections of pimonidazole hydrochloride (60-mg/kg body weight); 1 hour later they were killed and enucleated. The globes were fixed in 10% NBF for 2 hours; retinas were dissected and washed with tris-buffered saline (TBS); then they were blocked/permeabilized in 10% donkey serum with 1% Triton X-100/0.05% Tween 20 in TBS for 6 hours and stained with an antibody against pimonidazole-adducts (Hypoxyprobe, Burlington, MA, USA) followed by the secondary anti-rabbit IgG conjugated to Alexa Fluor 647- and Alexa Fluor 488-conjugated isoclin B4. The retinas were mounted on microscope slides with Prolong Gold mounting medium (Life Technologies). Ex vivo images were captured using an epifluorescence Nikon Eclipse Ti-E inverted microscope. Quantitative estimation of hypoxia in the RVO retinas were assessed using magic wand tool in Adobe image processing software (San Jose, CA, USA).

**Ultra-High Resolution Optical Coherence Tomography Imaging**

Optical coherence tomography of the retinas was performed using Bioptigen ultra-high resolution spectral-domain OCT (SD-OCT) imaging system with a mouse retinal bore (Bioptigen LLC, Morrisville, NC, USA). Before and at 2 hours after PRVT,
retinas of RVO mice were visualized with the SD-OCT system. Mice were anesthetized by intraperitoneal injection of a ketamine/xylazine (25/10 mg/kg) mixture and a 1% tropicamide solution was used to dilate the eyes for the retinal imaging. Genteal lubricating eye gel was used to keep the eyes moist. The mice were wrapped in gauze, placed in a holding chamber, and the head position was stabilized with a bite bar. Images were acquired from the optic disc to approximately 1.4 mm of the superior retina including the occlusion. InVivoVue software (Bioptigen LLC) was used for morphometric analysis of the retina.

Fluorescein Angiography

Fluorescein angiography was performed according to the methods previously described.31 Briefly, mice were anesthetized with a ketamine/xylazine (25/10 mg/kg) mixture; dilated with 1% tropicamide, and placed on a warm platform; Genteal lubricating eye gel was used to keep the eyes moist. A 10% sodium fluorescein (AK-fluor; Akorn, Decatur, IL, USA) sterile solution was injected systemically at a dose of 60 mg/kg; fluorescent and bright-field fundus images and in some cases videos were acquired using the Micron IV retinal imaging system.

Statistics

Data are presented as mean ± SD. Student’s t-tests were performed to compare two samples and, for comparison of more than two samples, 1-way ANOVA was performed using Prism 6 (Graph-Pad, San Diego, CA, USA). P less than or equal to 0.05 was considered as statistically significant.

RESULTS

Retinal Vein Occlusion Causes Acute Retinal Hypoxia at an Early Stage

To evaluate retinal hypoxia in RVO mice, one or two of the major branch retinal vein(s) was/were occluded by PRVT.29 Retinal hypoxia was detected in mouse RVO using the standard ex vivo pimonidazole-adduct immunostaining technique.28 Total occlusion of the retinal vein was confirmed by FA at 2 hours after the laser treatment (Fig. 1). Retinal hypoxia was observed in all RVO mice within 2 hours post-PRVT. Photodynamic retinal venous thrombosis induced occlusion of one major retinal vein resulted in patchy areas of extravascular hypoxia adjacent to the occluded vein in the peripheral retina. Due to the dichotomous branching of the occluded retinal vein, regions of hypoxia were observed throughout one-half of the retina. Based on pimonidazole-adduct immunostaining, we estimate that approximately 12% and 30% of the mouse retina becomes hypoxic following one or two vaso-occlusive events (nasal and temporal), respectively (Fig. 3). There were no significant changes in the retinal hypoxia of the nasal vessels after temporal vein occlusion. In transverse retinal sections from these RVO mice, hypoxia was localized to the cells in the inner nuclear layer with processes extending to the inner plexiform layer and retinal ganglion cell layer (Supplementary Fig. S4). After PRVT, the venous diameter downstream to the occlusion site is threadlike and is barely appreciable by FA (Fig. 1). Also, upstream from the occlusion site, the vein becomes dilated as seen in the fluorescein angiography. Supplementary Movie S1 illustrates an example of the venous blood circulation after RVO induction, with blood flow redirected toward the central retina presumably through collateral circulation,52 as evidenced by increased fluorescence observed in FA.

Retinal Vein Occlusion Leads to Retinal Edema, Increased Vascular Tortuosity and Neovascularization

Retinal vein occlusion causes focal retinal tissue damage that was clearly visible in the b-scan and en face images obtained from an SD-OCT imaging system (Supplementary Fig. S1). We observed changes in retinal thickness shortly after PRVT-
induced occlusion that most likely result from increased permeability of the occluded vein. This proposition is supported because we applied the same amount of energy via PRVT to retinal areas located between two major vessels without significantly affecting retinal thickness. We also examined the retinal thickness at later times post-PRVT revealing that the edema observed earlier, had resolved to normal levels by day 8 (Supplementary Fig. S2). Retinal hemorrhages and increased vascular tortuosity were also commonly observed shortly after the vaso-occlusive event. Similar to other studies, in these RVO mice, the occluded vein spontaneously recanalized approximately 1 week post-PRVT treatment, as demonstrated by FA. Interestingly, nominal NV was commonly observed in these mice approximately 10 to 14 days post-PRVT (Fig. 4; Supplementary Fig. S3).

In Vivo and Ex Vivo Detection of Retinal Hypoxia in RVO Using HYPOX-4

In vivo imaging of retinal hypoxia was performed in C57BL/6 RVO mice using HYPOX-4. HYPOX-4 was systemically administered 2 hours after the PRVT; 22 hours later, HYPOX-4–dependent fluorescence was detected, indicating hypoxia upstream from the photocoagulation site (Fig. 5). Ex vivo analysis of HYPOX-4 fluorescence from the same retinal tissues, as measured by computer aided analysis (ImageJ software, http://imagej.nih.gov/ij/; provided in the public domain by the National Institutes of Health, Bethesda, MD, USA) agreed with the in vivo findings. The patterns of retinal hypoxia observed by HYPOX-4–dependent imaging were generally the same as those obtained by pimonidazole-adduct immunostaining. HYPOX-4 produced more confluent images of retinal hypoxia, whereas the pimonidazole-adduct staining patterns appeared patchy. We did not observe any change in autofluorescence in mouse retinal tissues due to PRVT-induced RVO (Supplementary Fig. S5). Also, HYPOX-4 was cleared from ocular tissues within 24 hours after injection (Supplementary Fig. S6). These results clearly demonstrate that vascular occlusion can induce retinal hypoxia, in addition to demon-
strating the utility of HYPOX-4 to detect it in the in vivo setting. The results suggest that in vivo HYPOX-4–dependent retinal imaging could provide useful information about the role of hypoxia in RVO.

**DISCUSSION**

Increased ocular VEGF has been reported in the mouse and macaque in response to PRVT-induced RVO. Because VEGF is hypoxia-dependent, and ischemia is present in these models, it is reasonable to assume that it is an indicator of retinal hypoxia. Oxygen-electrode measurements in miniature pigs with PRVT-induced RVO, revealed preretinal hypoxia over the occluded vein, thus providing a direct measure of retinal hypoxia due to RVO. Similar to pimonidazole, the structure of HYPOX-4 incorporates the nitroimidazole moiety, serving as a biomarker of tissue hypoxia. In tissues with low oxygen pressure, azo- and nitroreductases are hypothesized to reduce the nitroimidazole causing it to become protein bound and sequestered. Using standard optical imaging equipment, Oregon Green dye also incorporated into the HYPOX-4 structure, enables the detection of zones of retinal hypoxia by its fluorescence activity. We have demonstrated here, through the application of HYPOX-4, that retinal hypoxia develops after the onset of RVO-induced retinal ischemia in living mice and in real time. Heretofore, these types of measurements have not been reported in animal models of RVO; which may be largely due to the limitations of existing technologies. The observed patterns of retinal hypoxia determined by HYPOX-4–dependent fluorescence, are consistent with those that would be expected, overlapping significantly with the retinal areas that are drained by the vein prior to the occlusive event (Fig. 5). Our data show that retinal hypoxia increases with the number of PRVT-induced occlusions. Patterns of hypoxia determined by HYPOX-4 fluorescence activity were generally similar to those obtained by the ex vivo pimonidazole-adduct immunofluorescence technique. Although, they tended to be nominally more diffuse, probably owing to the greater bioavailability of HYPOX-4. The pimonidazole immunoreactivity provided an alternative hypoxia assessment and confirmed HYPOX-4 as a potential in vivo hypoxia-imaging agent.

We observed retinal edema and NV in our RVO mice; both are vision-threatening sequelae to RVO. In patients, retinal edema may produce a serious distortion of vision if the macula is involved. It has been presumed that venous occlusion results in increased fluid pressure, causing the affected vein to swell, compromising its structural integrity, allowing fluid to leak into the retinal parenchyma, and finally resulting in macular edema. Notably, clinical and experimental data demonstrate the efficacy of anti-VEGF therapies against macular edema secondary to RVO. This immediately raises the suspicion of hypoxia as a pathogenic component, although other molecular signaling mechanisms that raise tissue levels of VEGF are known. Interestingly, if hypoxia does indeed factor into the pathogenesis of RVO-associated macular edema, then its in vivo imaging would offer distinct advantages. In this study, we observed retinal edema 2 hours post-PRVT that resolved by day 8 (Supplementary Fig. S2). And this finding is consistent with the results of other studies in the mouse, rat, and pig (Supplementary Fig. S1). At this early time-point, the observed retinal edema is most likely due to increased permeability of the distended and structurally compromised occluded retinal vein, rather than hypoxia. This is reasonable because this time of assessment is too short for de novo protein synthesis to occur (e.g., hypoxia-induced VEGF or other vasoactive factors). We observed retinal NV in all of the eyes of these mice 10 to 14 days post-PRVT (Fig. 4; Supplementary Fig. S3). These results are generally consistent with those of previous studies; retinal NV, secondary to RVO, was observed at 2 weeks in the mouse, 3 weeks in miniature pigs, 2
weeks in albino rats, and 10 to 12 weeks in pigmented rats.

In each of these studies, recannulization of occluded retinal veins in experimental subjects was reported. We observed recannulization of the affected retinal vein by post-PRVT day 8, in almost all of our RVO mice. Recannulization clearly increases blood flow, oxygenating the retina, and conceivably reducing any hypoxia-dependent VEGF. Therefore, it is reasonable to consider whether the retinal NV that others and we have observed is indeed hypoxia-dependent. After laser-induced venous occlusion, blood flow and oxygen diffusion to the perivascular tissue ceases, and by our HYPOX-4 assessments, triggering the onset of hypoxia. Additionally, destructive processes occurring secondary to venous occlusion may ensue, creating focal regions of retinal hypoxia due to vasoregression. This may offset the oxygenation effects of recannulization, resulting in a hypoxia-induced vasoproliferative response. In the current study, we assessed retinal hypoxia at 9 (HYPOX-4 in vivo) to 10 days (Pimonidazole ex vivo) prior to the development of retinal NV. Our data do not support or refute a link between hypoxia and retinal NV in this mouse model of RVO. Furthermore, we do not discount the possibility that ischemia-reperfusion injury or inflammation may raise VEGF levels contributing to the observed retinal NV. Future studies are aimed at better defining the hypoxia/NV relationship in experimental RVO using HYPOX-4 as a molecular probe to detect and quantify retinal hypoxia. The development of a durable RVO model in rodents will be critical to these studies and we are currently pursuing this goal.

In conclusion, we have demonstrated that retinal hypoxia and edema appear within 2 hours, and retinal NV at 10 to 14 days post-PRVT in RVO mice. To our knowledge, this is the first report describing the occurrence of retinal hypoxia and edema at this early time-point. This study also demonstrates that, HYPOX-4, a new optical imaging probe, has high potential for the detection of retinal hypoxia in vivo in experimental RVO, before the onset of physiological changes leading to retinal cell damage and NV. We propose that the level of retinal hypoxia produced by a PRVT-induced occlusive event in the retina is proportional to the intensity of a developing vasoproliferative response. In future studies, we will test the ability of HYPOX-4–dependent imaging of hypoxia to determine this relationship and hopefully use it to predict the potential severity of an incipient vasculopathy.

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**Figure 5.** HYPOX-4–dependent in vivo and ex vivo imaging of retinal hypoxia in a RVO mouse. HYPOX-4 was administered by intraperitoneal injection 2 hours post-PRVT. In vivo imaging was performed 22 hours post injection. (A) Bright-field fundus photograph of a RVO mouse. The arrowhead and yellow circle show the position of the vein occlusion. (B) In vivo HYPOX-4–dependent fluorescence activity was detected upstream from the occlusion, indicating retinal hypoxia (green). (C) Ex vivo HYPOX-4–dependent fluorescence activity detected in the same retina (green).

(D) Relative fluorescence intensity was measured using ImageJ software from hypoxic and normoxic retina in (B) and represented as RFU. (E) Quantitative analysis of HYPOX-4–dependent fluorescence in RVO retinas receiving single or double vein occlusion. Student’s t-test was performed and P < 0.001 is considered as highly significant. ONH, optic nerve head.
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References


