Ferrochelatase is a therapeutic target for ocular neovascularization

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Abstract

Ocular neovascularization underlies major blinding eye diseases such as “wet” age-related macular degeneration (AMD). Despite the successes of treatments targeting the vascular endothelial growth factor (VEGF) pathway, resistant and refractory patient populations necessitate discovery of new therapeutic targets. Using a forward chemical genetic approach, we identified the heme synthesis enzyme ferrochelatase (FECH) as necessary for angiogenesis in vitro and in vivo. FECH is overexpressed in wet AMD eyes and murine choroidal neovascularization; siRNA knockdown of Fech or partial loss of enzymatic function in the Fech<sup>−/−</sup> Pas<sup>−/−</sup> mouse model reduces choroidal neovascularization. FECH depletion modulates endothelial nitric oxide synthase function and VEGF receptor 2 levels. FECH is inhibited by the oral antifungal drug griseofulvin, and this compound ameliorates choroidal neovascularization in mice when delivered intravitreally or orally. Thus, FECH inhibition could be used therapeutically to block ocular neovascularization.

Keywords  age-related macular degeneration; angiogenesis; ferrochelatase; griseofulvin; heme synthesis

Subject Categories  Neuroscience; Pharmacology & Drug Discovery; Vascular Biology & Angiogenesis

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Introduction

Ocular neovascularization is a characteristic of blinding eye diseases like retinopathy of prematurity, proliferative diabetic retinopathy, and “wet” age-related macular degeneration (AMD; Das & McGuire, 2003). These diseases cause vision loss throughout life and together are the major cause of blindness in developed countries (Das & McGuire, 2003; Penn et al, 2008). All are characterized by abnormal angiogenesis (neovascularization) in the posterior segment of the eye (in the retina or choroid), leading to vascular leakage that can result in hemorrhage, edema, and retinal detachment, with vision loss as a direct result. This neovascularization is driven by a number of factors, prominent among which is vascular endothelial growth factor (VEGF; Penn et al, 2008).

Management of neovascular eye diseases has been revolutionized by the advent of anti-VEGF biologics such as ranibizumab, bevacizumab, and aflibercept (Folk & Stone, 2010; Prasad et al, 2010; Hanout et al, 2013). However, despite these considerable advances, treatment options remain limited for a substantial fraction of patients who are non-responsive or refractory to anti-VEGF therapy (Lux et al, 2007); no agents acting downstream of the VEGF receptor interaction are yet approved for targeted therapy of these diseases (Abouammoh & Sharma, 2011; Lally et al, 2012). There is thus a pressing need to identify alternative targets in angiogenesis signaling that could form the basis for new therapeutics. To uncover novel, potentially druggable angiogenesis mediators, here we took a forward chemical genetic approach to find protein targets of an antiangiogenic natural product, cremastranone (Shim et al, 2004; Kim et al, 2007, 2008; Lee et al, 2014; Basavarajappa et al, 2015). We identify ferrochelatase (FECH) as a target of this compound and show that it is necessary for angiogenesis in vitro and in vivo and targetable with the FDA-approved small molecule drug, griseofulvin. FECH is the terminal enzyme in heme biosynthesis, responsible for catalyzing the insertion of Fe<sup>2+</sup> ion into protoporphyrin IX (PPIX; Hamza & Dailey, 2012). To our knowledge, we describe here for the first time FECH’s role as a druggable mediator of angiogenesis.
Results

Ferrochelatase is a target of an antiangiogenic compound

To identify potentially novel protein modulators of angiogenesis, we used photoaffinity chromatography to search for targets of the naturally occurring antiangiogenic compound, cremastranone (Fig 1A), which has selective antiproliferative effects on endothelial cells (Lee et al., 2014). Protein binding partners of cremastranone were pulled down from a tissue lysate using immobilized cremastranone-based affinity reagent but not a negative control reagent (Fig 1B). One of the two pulled-down proteins was identified using peptide mass fingerprinting as FECH (Fig 1C and Appendix Fig S1). Immunoblot of eluates from photoaffinity pull-down experiments confirmed the identity of the pulled-down protein using an antibody against FECH (Fig 1D). In order to confirm specificity of binding between cremastranone and pulled-down proteins, the affinity reagent was incubated with tissue proteins in the presence of excess active cremastranone isomer (Fig 1B; Basavarajappa et al., 2014). Under this condition, the binding of FECH to the affinity reagent was markedly decreased (~87%), indicating competition for binding to FECH between the cremastranone isomer and the affinity reagent (Fig 1E).

Recombinant FECH also interacted with the affinity reagent (Fig 1F and G), indicating that the interaction does not require eukaryotic accessory proteins. Some binding to the negative control is likely due to the abundance of recombinant protein present; recent work confirms that cellular FECH does not readily bind linkers such as our control compound (Park et al., 2016). Moreover, cremastranone treatment of human retinal endothelial cells (HRECs) caused a dose-dependent buildup of the FECH substrate PPIX (Fig EV1A), indicative of FECH inhibition, and addition of excess 5-aminolevulinic acid (5-ALA; the first precursor compound in the heme biosynthetic pathway that promotes increased heme production) partially rescued cremastranone’s antiproliferative effects on HRECs (Fig EV1B). Cremastranone did not chelate iron, suggesting that it does not act indirectly on FECH by sequestering FECH’s Fe2+ substrate (Fig EV1C). The FECH pathway, therefore, is targeted by a known antiangiogenic compound, suggesting that this protein and pathway are important for angiogenesis.

Figure 1. Ferrochelatase (FECH) is a target of the antiangiogenic natural product, cremastranone.
A Chemical structure of cremastranone (1) (top). Antiproliferative activity of cremastranone (bottom) shown as the 50% growth inhibitory concentration (GI50) on human retinal endothelial cells (HRECs), human umbilical vein endothelial cells (HUVECs), and non-endothelial ocular cells (uveal melanoma 92-1, retinoblastoma Y-79, and retinal pigment epithelium ARPE-19).
B Structures of compounds used in photoaffinity chromatography.
C Proteins pulled down with indicated reagents (numbered as in B) in photoaffinity chromatography were separated on SDS–PAGE and silver-stained.
D Immunoblot of pulled-down proteins using antibody against ferrochelatase (FECH).
E Immunoblot of pulled-down proteins from competition assay with excess active cremastranone isomer (4), relative quantification of band intensity shown.
F Silver-stained SDS–PAGE gel of recombinant human FECH protein pulled down using photoaffinity chromatography.
G Anti-FECH immunoblot of recombinant human FECH protein pulled down using photoaffinity chromatography.

Data information: All the gel and immunoblot images are representative from at least two independent experiments. Source data are available online for this figure.
**Ferrochelatase is required for angiogenesis in vitro**

The role of FECH in angiogenesis has not previously been explored, despite strong evidence linking heme catabolism with angiogenesis (Dulak et al, 2008). To determine whether FECH plays a key role in angiogenesis, we knocked down FECH using siRNA (Fig 2A and B) and monitored key angiogenic properties of HRECs in vitro. FECH knockdown significantly reduced the proliferation of HRECs (Figs 2C and EV2A). There was also a significant decrease in migration of HRECs treated with FECH siRNA in a scratch-wound assay (Fig 2D). Further, knocking down FECH in HRECs completely abolished the tube formation ability of HRECs as monitored in a Matrigel assay (Fig 2E).

To confirm the knockdown results, we also tested whether a known pharmacological inhibitor of FECH shows antiangiogenic properties in vitro. N-methyl protoporphyrin (NMPP), a competitive inhibitor of FECH activity (Cole & Marks, 1984), also inhibited proliferation, migration, and tube formation ability of HRECs in vitro (Fig 2F–H). Similar results were seen in the primate choroidal endothelial cell line, Rf/6a (Fig EV3A and B), broadening the relevance of these findings. However, despite these potent antiproliferative effects, FECH knockdown and low-dose chemical inhibition were not associated with significant apoptosis of these cells (Fig EV2B–D), indicating a cytostatic rather than cytotoxic effect. This was further confirmed in a washout experiment: HRECs resumed proliferation after removal of NMPP (Fig EV2E). Moreover, FECH knockdown did not inhibit proliferation of non-endothelial ocular cell lines ARPE-19 and 92-1 as well as macrovascular human umbilical vein endothelial cells (HUVECs; Fig EV4A–C), indicating that FECH inhibition is not associated with general cytotoxicity. Likewise, NMPP had negligible antiproliferative effect on these cells and on retinoblastoma cell line Y-79 (Fig EV4D–F and J). However, NMPP had similar antiproliferative potency on human brain microvascular endothelial cells (BMECs) to its effects on HRECs (Fig EV4L). Together, these experiments confirm that FECH function is required for angiogenesis in vitro and that the angiostatic effects of FECH blockade are not cytotoxic and may be microvascular cell-specific.

**Ferrochelatase is upregulated during neovascularization**

Given the potent antiangiogenic effects of FECH knockdown we observed in culture, we then explored whether FECH is associated with neovascularization in vivo. We employed a mouse model of laser-induced choroidal neovascularization (L-CNV). This widely used model recapitulates some of the features of wet AMD (Montezuma et al, 2009). FECH was overexpressed in and around lesions during neovascularization in this model (Fig 3A); this overexpression was seen throughout the retina, but especially in neovessels (Fig 3B). More importantly, FECH expression was seen throughout the retinal layers of human wet AMD patients analyzed postmortem. In the subretinal layers including the choroid (the origin of neovascularization in wet AMD), expression was significantly increased compared to healthy, age-matched eyes (Fig 3C).

**Ferrochelatase is required for neovascularization in vivo**

Since FECH upregulation suggested a role for this protein in neovascularization in the living eye, we asked whether decreased FECH would inhibit this process. When L-CNV mice were treated intravitreally with Fech-specific siRNA, there was a significant decrease in choroidal neovascularization as compared with both saline-treated control mice and control non-targeting siRNA-treated mice (Fig 3D). To confirm these findings in a genetic model, we turned to the Fech<sup>m1Pas</sup> mouse, which carries a partial loss-of-function M98K point mutation in the Fech gene (Tutuo et al, 1991). In M98K heterozygotes, FECH activity is 45–65% of normal, while in homozygotes, activity is < 10% of normal (Boulechar et al, 1993). We observed a reduction in L-CNV in heterozygous Fech<sup>m1Pas</sup> mutant mice compared with wild type, and this reduction was even more pronounced in the Fech<sup>m1Pas</sup> homozygotes (Fig 3E). These in vivo experiments confirm the clinical relevance of FECH in neovascularization, and the value of targeting this enzyme to block this process.

**Ferrochelatase-targeting therapy treats neovascularization**

The FDA-approved antifungal drug, griseofulvin, has been in clinical use for over half a century (Petersen et al, 2014). The primary antifungal mechanism of this compound is as a microtubule inhibitor (Borgers, 1980). However, an unexpected off-target effect of this drug is inhibition of FECH (Holley et al, 1990; Brady & Lock, 1992; Martinez et al, 2009). Griseofulvin alkylates the heme prosthetic group of cytochrome P450 in vivo, forming NMPP, the FECH active-site inhibitor (Liu et al, 2015). Taking advantage of this property, we treated HRECs with griseofulvin and observed dose-dependent antiproliferative effects, inhibition of migration, and inhibition of tube formation (Fig 4A–C) similar to those observed after FECH knockdown. The concentrations of griseofulvin needed to have effects on endothelial cells were higher than those seen with NMPP (Fig 2F–H), likely due to incomplete alkylation of heme in griseofulvin-treated cells. However, this concentration (~10 μM or ~3.5 ng/ml) is 2.75 logs less than that attained in plasma during antifungal treatment of humans (1–2 μg/ml; Epstein et al, 1972), suggesting that efficacy in vitro can be achieved in a clinically attainable concentration range. Moreover, effective antiangiogenic concentrations of griseofulvin were not associated with apoptosis (Fig EV2F) of HRECs and could be washed out with resumption of proliferation (Fig EV2G). Griseofulvin did not have pronounced antiproliferative effects on other ocular cell types or HUVECs (Fig EV4G–I and K), but did affect proliferation and tube formation of Rf/6a choroidal endothelial cells (Fig EV3C and D) and proliferation of BMECs (Fig EV4M). In addition, griseofulvin inhibited formation of microvascular sprouts in the choroidal sprouting assay, an <i>ex vivo</i> model of choroidal angiogenesis (Fig 4D). It also performed similarly in an <i>ex vivo</i> retina endothelial sprouting assay, showing pronounced antiangiogenic activity (Fig 4E). Thus, griseofulvin specifically inhibits the growth of microvascular endothelial cells, including those of the retina and choroid.

Spurred by these findings, we tested griseofulvin as a therapy for L-CNV. <i>Ad libitum</i> feeding of ≥ 0.5% (w/w) griseofulvin to mice has previously been demonstrated to induce inhibition of FECH (Brady & Lock, 1992). Excitingly, such oral treatment of L-CNV mice decreased neovascularization (Figs 5A–C and EV5A and B); similarly, dose-dependent results were seen when griseofulvin was intravitreally injected instead; this is the standard delivery route for existing anti-VEGF agents (Fig 5D–F). Both delivery routes had
Figure 2. FECH is an essential protein for angiogenesis in vitro.

A FECH is knocked down using specific siRNAs as confirmed by immunoblot.
B Quantification of FECH knockdown. *P = 0.0319, two-tailed unpaired Student’s t-test (n = 3 per group).
C Proliferation of HRECs was monitored in presence or absence of FECH-specific siRNA. **P = 0.0004, two-tailed unpaired Student’s t-test (n = 6 per group).
D Scratch-wound migration assay was performed with or without FECH knockdown in HRECs. ***P = 0.0007, two-tailed unpaired Student’s t-test (n = 3 per group).
E Ability of HRECs to form tubes in vitro on Matrigel was monitored after knocking down FECH. ***P = 0.0003, two-tailed unpaired Student’s t-test (n = 6 per group).
F The effect of NMPP, a specific inhibitor of FECH activity, on in vitro proliferation was measured using an AlamarBlue assay (n = 3 per dose).
G Migration of HRECs in presence of different doses of NMPP was measured using a scratch-wound assay. ****P = 0.0001 compared to DMSO-treated sample, ANOVA with Dunnett’s post hoc tests (n = 3 per group). Dotted lines indicate borders of scratch at time zero.
H Ability of HRECs to form tubular structures in Matrigel was monitored after NMPP treatment. ****P = 0.0001 compared to DMSO-treated sample, ANOVA with Dunnett’s post hoc tests (n = 4 per group).

Data information: Graphs show mean ± SEM. Representative results from at least three independent experiments. Scale bars = 1 mm.
Source data are available online for this figure.
similar maximal effects. We also tested whether intravitreal griseofulvin could synergize with anti-VEGF164 antibody treatment, equivalent to the standard of care. We observed a modest additive effect of low-dose anti-VEGF164 with 50 μM griseofulvin (Fig EV5C), perhaps warranting further exploration of this combination therapy.

**Ferrochelatase depletion decreases the VEGF receptor via eNOS**

In order to understand how FECH contributes to angiogenesis, we knocked down FECH in HRECs and monitored levels of endothelial nitric oxide synthase (eNOS), a key angiogenesis regulator that

Figure 3. FECH is an essential protein for angiogenesis in vivo.
A Whole mount staining of L-CNV mouse choroid stained with an antibody against FECH (red) and with agglutinin (green) to label neovascularization. Scale bar = 50 μm.
B Sections of L-CNV and untreated mouse eyes stained with an antibody against FECH (red), with agglutinin (green) to label neovascularization and with DAPI to label nuclei (blue). Retinal layers indicated: GCL, ganglion cell layer; INL, inner nuclear layer; ONL, outer nuclear layer. Scale bar = 20 μm.
C Immunostaining of sections of eyes from an 89-year-old wet AMD patient and age-matched control using an antibody against FECH (red). The nuclei of cells are stained blue with DAPI. Retinal layers indicated: GCL, ganglion cell layer; INL, inner nuclear layer; ONL, outer nuclear layer. Scale bar = 20 μm. The staining intensity of subretinal FECH, where CNV occurs, was quantified in human subjects (n = 3 per group). *P = 0.0269, two-tailed unpaired Student’s t-test.
D Whole mount staining of RPE/choroid isolated from L-CNV mice treated with Fech-specific siRNA. The choroidal vasculature was stained with agglutinin conjugated with Alexa Fluor 555 (red), and the neovascular area around the lesions was quantified. *P = 0.044; **P = 0.0038, ANOVA with Tukey’s post hoc tests (n = 5–9 per group). Scale bar = 50 μm.
E Whole mount staining of RPE/choroid isolated from mice wild-type (WT), heterozygous (Hetero), or homozygous (Homo) for the Fech<sup>11<sup>th</sup></sup> point mutation that underwent L-CNV. The choroidal vasculature was stained with agglutinin conjugated with Alexa Fluor 555 (red), and the neovascular area around the lesions was quantified. *P = 0.015; **P = 0.0037 versus WT, ANOVA with Tukey’s post hoc tests (n = 9–13 per group). Scale bar = 50 μm.

Data information: Graphs show mean ± SEM.
Figure 4. Chemical inhibition of FECH inhibits angiogenesis in vitro.

A The effect of griseofulvin, an FDA-approved drug that inhibits FECH activity, on proliferation of HRECs in vitro. (n = 3 per dose).
B Migration of HRECs in presence of griseofulvin was measured in a scratch-wound assay. ****P = 0.0001, ANOVA with Dunnett’s post hoc tests (n = 3 per group).
C Ability of HRECs treated with griseofulvin to form tubular structures in Matrigel was monitored and analyzed using ImageJ. **P = 0.0082, ANOVA with Dunnett’s post hoc tests (n = 6 per group).
D The mouse choroidal sprouting assay was used to further measure the antiangiogenic potential of griseofulvin. Representative phase contrast images and quantitative analysis of choroidal endothelial sprouting. **P = 0.0018; ****P = 0.0001 versus DMSO at same time point, two-way repeated measures ANOVA with Dunnett’s post hoc tests (n = 3–4 per group).
E The ex vivo murine retinal angiogenesis assay was also used to measure the antiangiogenic potential of griseofulvin. Representative phase contrast images and quantitative analysis of retinal endothelial sprouting. ****P = 0.0001 versus DMSO at same time point, two-way repeated measures ANOVA with Dunnett’s post hoc tests (n = 5 per group).

Data information: Graphs show mean ± SEM, and the images are representative results from at least two independent experiments. Scale bars = 1 mm.
requires a heme cofactor for its activity to produce the proangiogenic molecule nitric oxide (NO; Förstermann & Münzel, 2006). Immunoblot of HRECs treated with FECH siRNA showed significantly decreased levels of eNOS (Fig 6A). Further, NOS activity was decreased in FECH knockdown HRECs as compared to negative control siRNA-treated cells (Fig 6B). To assess hemylation of eNOS,
we used a hemin pull-down assay. Only hemoproteins in the apo form (enzyme lacking the heme cofactor) will bind to hemin agarose beads. Despite the overall decrease in eNOS levels, we observed that the remaining eNOS was in the apo form (Fig 6C) when HRECs were treated with NMPP, the specific inhibitor of FECH enzymatic activity. As NO, the product of eNOS enzymatic activity, stabilizes hypoxia inducible factor (HIF) 1α (Sandau et al, 2001), we also monitored levels of this key mediator of angiogenesis after knocking down FECH in HRECs. 

**FECH knockdown decreased protein levels HIF-1α (Fig 6D).**

Since VEGF is a major proangiogenic stimulus, we monitored key events of VEGF signaling after knocking down FECH. We observed a profound decrease in (activating) phosphorylation of the major VEGF receptor VEGFR2 as well as total VEGFR2 protein levels in FECH knockdown cells when cultured in basal medium containing VEGF165 (Fig 6E), although production of VEGFA mRNA was unchanged (Appendix Fig S2). Interestingly, levels of related receptors VEGFR1 and neuropilin 1 were not affected by FECH depletion (Fig 6E). The protein levels of eNOS and VEGFR2 were partially rescued after treating FECH knockdown cells with exogenously added hemin (a stable form of heme, the enzymatic product of FECH; Fig 6F). The decreased levels of eNOS, VEGFR2, and HIF-1α after FECH knockdown were not due to general translational inhibition: There was no difference in phosphorylation levels of eIF-2α in FECH siRNA-treated cells (Fig 6F).

### Discussion

Understanding the molecular events of pathological angiogenesis is key to developing novel therapeutic options for neovascular eye diseases such as proliferative diabetic retinopathy, retinopathy of prematurity, and wet AMD. Currently, the drug pipeline for these diseases is dominated by anti-VEGF biologics (Kaiser, 2013). Even though these drugs have been successfully used to halt the progression of disease in wet AMD and proliferative diabetic retinopathy patients, there is a significant patient population (~30%) who are resistant to these treatments (Folk & Stone, 2010; Prasad et al, 2010). Hence diversification of the drug pipeline with novel therapeutic agents with different mechanisms of action is required (Smith & Kaiser, 2014). Toward this end, we used a forward chemical genetics approach using an antiangiogenic natural product, cremastinone, to uncover new drug targets for angiogenesis.

Using a series of biochemical, in vitro, and in vivo studies, we have determined that FECH is a key protein involved in ocular angiogenesis that can be exploited for developing novel antiangiogenic drugs. A photoaffinity-based chromatographic technique identified FECH as a protein binding partner of cremastinone. Although, like other natural products, cremastinone likely exhibits polypharmacology (Sulaiman et al, 2014), our data indicate that it exerts its antiangiogenic activity at least partially through inhibition of FECH activity.

Interestingly, FECH is a bona fide mediator of angiogenesis. Lack of FECH activity caused inhibition of angiogenesis both in vitro and in vivo. More importantly, only retinal and choroidal microvascular endothelial cell proliferation was inhibited in vitro while other ocular cell types tested did not show significant decreases in cell proliferation after FECH knockdown. Even macrovascular HUVECs were not as profoundly affected by FECH depletion as the microvascular HRECs and Rβ/6α cells. These data reveal that microvascular endothelial cells are particularly sensitive to FECH depletion. The lack of cytotoxic effects of FECH inhibition leads to consideration of FECH as a novel therapeutic target for ocular neovascular disease, possibly with minimal side effects. Supporting this latter assertion, in the genetic disease erythropoietic protoporphyria (EPP), FECH activity is markedly reduced, but EPP patients infrequently present severe symptoms apart from skin photosensitivity (Lecha et al, 2009). It may be possible to tune FECH activity to minimize side effects while maximizing antiangiogenic effects: Heterozygous Fech<sup>−/+</sup> mice do not display an EPP phenotype, despite reduced FECH function (Boulechar et al, 1993), but do show reduced choroidal neovascular response (this study). Neovascularization in human EPP patients has not been studied extensively, although a single case report describes an ocular phenotype of idiopathic optic neuropathy in an EPP patient, not conclusively related to the disease (Tsiboli et al, 2007).

Excitingly, we have shown that the FDA-approved antifungal drug and FECH inhibitor griseofulvin inhibited ocular angiogenesis in the L-CNV mouse model when administered orally. Griseofulvin has been used widely to treat fungal infections and is taken orally, often for months or years (Petersen et al, 2014; Liu et al, 2015). An off-target side effect of this therapy is that griseofulvin causes the formation of NMPP, along with other alkylated porphyrins, primarily in the liver (Liu et al, 2015). NMPP in turn acts as an active-site inhibitor of FECH (Cole & Marks, 1984). As with genetic reduction in function of FECH, apart from skin photosensitivitiy, no other major, common side effects are reported with systemic griseofulvin treatment of humans (Elewski & Tavakkol, 2005; Grover et al, 2012).

The fact that griseofulvin-fed mice showed decreased ocular neovascularization as compared to control mice is important as
currently there are no oral drugs available for the treatment of ocular neovascularization. The current approved drugs (anti-VEGF biologics) are all delivered intravitreally. If side effects were manageable, an oral therapy would be appealing. However, since griseofulvin was also effective when delivered intravitreally, combination intravitreal therapies of this drug with anti-VEGF biologics are also an interesting possibility to increase efficacy and decrease side effects of existing treatments. Moreover, griseofulvin offers the
promise of low-cost treatment if shown efficacious in clinical trials. There are also prospects for novel FECH inhibitors as therapy: We have developed cremasterona analogs that may also act through FECH (Basavarajappa et al., 2015), and recently, the drugs vemurafenib (Savitski et al., 2014) and salicylic acid (Gupta et al., 2013), as well as a lipid-derived probe compound (Niphakis et al., 2015), were all shown to bind to FECH, suggesting that this protein has many potential binding clefts and is therefore likely druggable with novel compounds.

FECH is the only known enzyme in humans to synthesize heme, and any loss of activity of this enzyme might result in intracellular heme deficiency (Dailey et al., 2000). Heme serves as either a prosthetic group for many enzymes or as a signaling molecule regulating expression of genes involved in cell differentiation, proliferation, and signal transduction (Mense & Zhang, 2006). FECH knockdown decreased protein levels of eNOS, HIF-1α and VEGFR2, key players promoting angiogenesis, in HRECs. As eNOS is the heme-dependent enzyme, FECH knockdown directly affects the stability and activity of eNOS. The product of the eNOS enzyme, NO, induces angiogenesis (Ziche & Morbidelli, 2000), partly by stabilizing HIF-1α under normoxia (Sandau et al., 2001). This explains the logic behind the decreased protein levels of HIF-1α in FECH knockdown cells with decreased eNOS activity. HIF-1α is a transcription factor that regulates the expression of a range of proangiogenic proteins. VEGFR2, the major tyrosine kinase receptor of VEGF165, is upregulated under hypoxia (Tuder et al., 1995; Takagi et al., 1996). The decreased levels of VEGFR2 in HRECs after FECH siRNA treatment can be linked to decreased HIF-1α. Taken together, low protein levels of VEGFR2 (and thus angiogenesis) in FECH-deficient HRECs can be directly attributed to loss of heme in cells and the resultant decrease in levels of eNOS enzyme. VEGFR2 depletion in turn further suppresses eNOS levels. The rescue of VEGFR2 and eNOS expression by addition of exogenous hemin, a stable form of heme, further supports this mechanism.

Our findings reveal a previously undocumented, central role of FECH in ocular angiogenesis. They provide a rationale for clinical testing of griseofulvin in neovascular eye disease. In addition, it will be valuable to develop novel, FECH-targeted therapies for treating the debilitating ocular diseases caused by neovascularization.

Materials and Methods

Materials

EBM-2 and IMDM growth media were purchased from Lonza (Walkersville, MD, USA). HRECs, BMECs, and Attachment Factor were purchased from Cell Systems (Kirkland, WA, USA). Clonetics® HUVECs were purchased from Lonza. All endothelial cells were used between passages 5 and 8. Endothelial growth medium (EGM-2) was prepared by mixing the contents of an EGM-2 “Bullet Kit” (Cat no. CC-4176) with endothelial basal medium (EBM; Lonza). Rl/6a cells were obtained from ATCC (Manassas, VA, USA) and grown in EEMEM. Y-79 cells were grown in RB medium (Basavarajappa et al., 2015), while 92-1 (Basavarajappa et al., 2015) and ARPE-19 (ATCC) cells were grown in RPMI and DMEM, respectively, supplemented with 10% FBS and 1% penicillin-streptomycin as described earlier (Basavarajappa et al., 2015); identity was confirmed by STR profiling and cells were monitored regularly for mycoplasma contamination. Click-it TUNEL Alexa Fluor 594 imaging assay kit (Cat no. C10246) was purchased from Molecular Probes (Eugene, OR, USA). Monoclonal antibodies against α-tubulin (DM1A) and β-actin (AC40), protoporphyrin IX, 5-aminolevulinic acid, hemin, hemin agarose beads, griseofulvin, and l-arginine were purchased from Sigma-Aldrich (St. Louis, MO, USA). N-methyl protoporphyrin (NMP) and the primary antibodies against FECH (A-3 and C20) were obtained from Santa Cruz (Santa Cruz, CA, USA). Antibodies against cleaved caspase 3 (SA1E), phospho-VEGFR2 (Tyk 1175) (19A10), phospho-ELF2α (Ser 51) (9721), ELF2α (9722), and eNOS (49G3) were from Cell Signaling (Danvers, MA, USA). Anti-HIF-1α (241809), VEGFR1 (AF321), and anti-VEGF164 (AF-493-NA) antibodies were purchased from R&D systems (Minneapolis, MN, USA). The antibody against neuropilin 1 (ab81321) was purchased from Abcam (Cambridge, MA, USA). Secondary antibodies were from Thermo Fisher Scientific (Pittsburgh, PA, USA). The Trizol, TaqMan probes, qPCR master mix, and 5-ethynyl-2'-deoxyuridine (EdU) incorporation assay kit were procured from Life Technologies (Carlsbad, CA, USA). The iScript reverse transcriptase was from Bio-Rad ( Hercules, CA, USA), AbD Serotec (Kidlington, UK) was the source of the alamarBlue, while BD Biosciences (San Jose, CA, USA) supplied the Matrigel. 4,5-Diaminofluorescein diacetate (DAF-2 diacetate) was purchased from Cayman Chemicals (Ann Arbor, MI, USA). ECL Primary Western blotting detection reagent was purchased from GE Healthcare (Buckinghamshire, UK).

Preparation of photoaffinity reagents

Synthesis and characterization of affinity reagents 2 and 3 was performed as described (Lee et al., 2016). Compound 2 contains a trimethoxy derivative of cremeastorenone that is more synthetically tractable, but which maintains good cellular potency and selectivity, both as homoisoflavonoid (Basavarajappa et al., 2015) and when incorporated into the affinity reagent (Lee et al., 2016). Compounds 1 and 4 were synthesized as described (Basavarajappa et al., 2014; Lee et al., 2014), with purity > 95%. For pull-downs, Neutravidin agarose beads (1 ml of 50% slurry) were washed three times in buffer A containing 25 mM Tris–HCl pH 7.4, 150 mM NaCl, 2.5 mM sodium pyrophosphate, 1 mM phenylmethylsulfonyl fluoride (PMSF), 0.1 mM sodium orthovanadate, 10 µg/ml aprotinin, and 10 µg/ml leupeptin. The beads were then incubated with 100 µM affinity or control reagents 2 or 3 overnight at 4°C with rotation. The beads were blocked using 1 mM biotin solution prepared in buffer A for 1 h followed by incubation with 1 mg/ml cytochrome c solution for 1 h at 4°C. The beads were then washed three times with buffer A and resuspended in 1 ml.

Photoaffinity pull-down experiments

Flash-frozen porcine brain (20 g) obtained from the Purdue-Indiana University School of Medicine Comparative Medicine Program was homogenized in 50 ml buffer A using a tissue homogenizer. Porcine brain was chosen as it is readily available and a rich source of protein. The homogenate was centrifuged at 2,000 g for 5 min. The supernatant (S1) was then homogenized by 50 strokes of a Dounce homogenizer followed by 10 min sonication with amplitude of 60% in cycles of 10 s sonication on and 40 s sonication off (Q125 from
Qsonica, Newtown, CT, USA). The lysate was then centrifuged at 11,000 g for 30 min. The resulting supernatant (S2) and pellet (P2) fractions were both collected. The P2 pellet was resuspended in buffer B: 1% Triton X-100 + buffer A and then homogenized by 25 strokes of a Dounce homogenizer and centrifuged at 11,000 g for 30 min; supernatant (S3) was collected. Both S2 and S3 supernatants were equally divided, and each fraction was incubated with 500 μl photoaffinity or control reagent conjugated to Neutravidin beads for 75 min at 4°C with shaking.

The beads were collected by centrifugation at 500 g for 5 min, then resuspended in 1 ml of buffer B and irradiated with 365 nm UV light (Mercury bulb H44GS100 from Sylvania in a Blak-Ray 100A long-wave UV lamp with output of 25 mW/cm² at sample distance) in a 60-mm Petri dish for 30 min at 4°C. The beads were then washed twice in buffer B, followed by three washes in high-salt buffer containing 25 mM Tris–HCl pH 7.4, 350 mM NaCl, 1% Triton X-100, and 1 mM PMSF. The beads were then washed again in salt-free buffer containing 25 mM Tris–HCl, 1% Triton X-100, and 1 mM PMSF. After 5-min incubation, the beads were collected and any residual liquid was removed using a Hamilton syringe. The Neutravidin beads were then heated in 300 μl of 3× SDS–PAGE gel loading dye containing 30 μl of 2-mercaptoethanol for 10 min at 70°C to release the bound proteins. After heating, the contents were allowed to cool and after a quick spin the eluate was collected using a Hamilton syringe. The eluates were then analyzed in 4–20% gradient SDS–PAGE, and the protein bands were visualized using silver staining (Corson et al., 2011). The protein bands pulled down specifically by photoaffinity reagent were excised from the silver-stained SDS–PAGE gel and analyzed by mass spectrometry (IUSM Proteomics Core). Using Sequest™ algorithms and the swine database (UniProt), the identities of the pulled-down proteins were confirmed (Appendix Fig S1).

For competition experiments, S2 and S3 supernatants were incubated with affinity reagent-Neutravidin beads in the presence of 1 mM crenastramone isomer SH-11052 (4) (Basavarajappa et al., 2014) and then processed as described above.

Recombinant FECH

Recombinant human protein FECH was purified as described previously (Dailey et al., 1994). Briefly, Escherichia coli JM109 cells transformed with plasmid pHDTF20 encoding recombinant human FECH were grown in Circlegrow medium containing 100 μg/ml ampicillin for 20 h at 30°C. The cells were harvested and resuspended in solubilization buffer (50 mM Tris-MOPS pH 8.0, 1% sodium deoxycholate, 100 mM KCl, and 1 mM PMSF). The cell suspension was sonicated and then ultracentrifuged at 45,000 g for 30 min. The supernatant was subjected to cobalt-affinity chromatography, and the column was washed with solubilization buffer containing 20 mM imidazole. The protein was eluted with 250 mM imidazole in solubilization buffer. The protein eluate was then dialyzed in solubilization buffer containing 10% glycerol before storage at 4°C. Recombinant protein (200 μg) was used in pull-down experiments as above.

Immunoblot assay

Immunoblots were performed as described previously (Basavarajappa et al., 2014). Briefly, cell lysates were prepared by incubating the cells for 10 min at 4°C in NP-40 lysis buffer (25 mM HEPES pH 7.6, 150 mM NaCl, 1% NP-40, 10% glycerol, 1 mM sodium orthovanadate, 10 mM NaF, 1 mM PMSF, 10 μg/ml aprotinin, 1 μM pepstatin, 1 μM leupeptin) and then centrifuged at 12,000 g for 15 min at 4°C. Supernatant was collected, and protein concentration was determined using a Bradford assay. Equal amounts of total protein (40 μg) from each sample were resolved by 10% SDS–PAGE and transferred onto PVDF membranes. Proteins were immunoblotted with antibodies against FECH (clone A-3; 1:1,000 dilution), α-tubulin (1:1,000), VEGFR1 (1:500), phospho-VEGFR2 (1:500), VEGFR2 (1:500), phospho-ef2α (1:500), ef2α (1:500), HIF-1α (1:500), neuropilin 1 (1:500), β-actin (1:1,000), or eNOS (1:500). All of the dilutions were made in Tris-buffered saline-0.05% Tween-20 buffer containing 2% bovine serum albumin (BSA).

siRNA knockdown of FECH in cells

Cells were grown in 6-well plates until 80% confluency was achieved. Lipofectamine RNAiMAX reagent (7.5 μl, Life Technologies) mixed with 30 pmol of siRNAs was added to each well according to the protocol recommended by the manufacturer. For FECH knockdown, 15 pmol each of two siRNAs (SASI_Hs01_00052189 and SASI_Hs01_00052190; Sigma) was used, and for negative control, MISSION® siRNA Universal negative control was used. Fresh EGM-2 medium was added to the plate 24 h after transfection, and cells were used 48 h after transfection for further experiments except for the proliferation time course, for which 24 h after transfection the cells were trypsinized and seeded in a 96-well plate.

Cell proliferation assay

Proliferation of cells was monitored as described before (Basavarajappa et al., 2014). Briefly, 2,500 cells (after FECH knockdown in some experiments) in 100 μl of growth medium were plated in each well of 96-well clear-bottom black plates and incubated for 24 h. Griseofulvin, NMPP, or DMSO vehicle (final DMSO concentration of 1%) was added, and the plates were incubated for 44 h in 100 μl media at 37°C and 5% CO₂. For washout experiments, 1,500 cells were initially plated, and drug-containing medium was removed after 48 h and cells incubated in fresh growth medium for a further 24 or 48 h. AlamarBlue reagent (11.1 μl) was added to each well of the plate and 4 h later fluorescence readings were taken at excitation and emission wavelengths of 560 nm and 590 nm, respectively, using a Synergy H1 plate reader (BioTek, Winooski, VT, USA). For dose–response experiments, GI₅₀ was calculated using GraphPad Prism v. 7.0.

Migration assay

The migration of HRECs was monitored as described before (Basavarajappa et al., 2014). Briefly, HRECs were grown until confluency in 6-well plates and then serum starved overnight in EBM-2 medium. Using a sterile 10-μl pipet tip, a scratch was introduced in each well and fresh EGM-2 medium containing DMSO or different concentrations of compounds was added to the wells. For knockdown experiments, the scratch was introduced 48 h after transfection and fresh medium was added to the wells. Photographs of the
wells were taken at different time points, and the number of migrated cells into the scratched area was manually counted.

**In vitro Matrigel tube formation assay**

The ability of HRECs and RI/6a cells to form tubes *in vitro* was monitored as described before (Basavarajappa et al., 2015). Briefly, cells were treated with the indicated concentrations of compounds or DMSO or siRNAs for 48 h and then 15,000 cells in 100 µl of growth medium containing siRNAs, DMSO, or compounds were added to each well of a 96-well plate that was pre-coated with 50 µl of Matrigel basement matrix. Photographs of each well at different time points were taken to measure the tube formation using the Angiogenesis Analyzer plugin in ImageJ software (v.1.48; http://image.bio.methds.free.fr/ImageJ/?Angiogenesis-Analyser-for-ImageJ.html).

**Animals**

All animal experiments were approved by the Indiana University School of Medicine Institutional Animal Care and Use Committee and followed the guidelines of the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Visual Research. Wild-type female C57BL/6J mice, 6–8 weeks of age or timed pregnancies, were purchased from Jackson Laboratory (Bar Harbor, ME) and housed under standard conditions (Wenzel et al., 2015). Foch TriPas mice (Tutois et al., 1991) were purchased from Jackson Laboratory on a BALB/c background and backcrossed into C57BL/6J; mixed-sex littermates from a subsequent brother–sister mating were used for experiments at 6–8 weeks of age. Mice were anesthetized for all procedures by intraperitoneal injections of 80 mg/kg ketamine hydrochloride and 10 mg/kg xylazine. Sample sizes were based on power analyses, and treatments were randomly assigned by cage.

**L-CNV model**

L-CNV was generated as described previously (Lambert et al., 2013; Poor et al., 2014; Sulaiman et al., 2015). Studies were powered to have an 80% chance of detecting effect size differences of 50%, assuming 30% variability, α = 0.05. Briefly, eyes were diluted using 1% tropicamide, then underwent laser treatment using 50 µm spot size, 50 ms duration, and 250 mW pulses of an ophthalmic argon green laser, wavelength 532 nm, coupled to a slit lamp. Where indicated, intravitreal injections of PBS vehicle, siRNA (1.25 µM final intravitreal concentration), griseofulvin (50, 100 µM, final intravitreal concentrations), anti-VEGF<sub>164</sub> antibody (0.2, 1, 5 ng delivered), or combinations of griseofulvin and anti-VEGF<sub>164</sub> antibody, as indicated, were given in a 0.5 µl volume at the time of laser treatment. Eyes were numbed with tetracaine solution before the injection, and triple antibiotic ointment was used immediately after the injection to prevent infection. A masked researcher undertook imaging and analysis to avoid bias. One week after laser treatment, mice underwent optical coherence tomography using a Micron III imager (Phoenix Research Labs, Pleasanton, CA, USA) and CNV lesions were quantified as ellipsoids as described (Sulaiman et al., 2015). Two weeks after laser treatment, eyes were enucleated and fixed, choroidal flat mounts prepared, and vasculature stained with rhodamine-labeled *Ricinus communis* agglutinin 1 (Vector Labs, Burlingame, CA, USA), followed by confocal Z-stack imaging (LSM 700, Zeiss, Thornwood, NY, USA) to estimate lesion volume. The sum of the stained area in each section, multiplied by the distance between sections (3 µm), gave the CNV lesion volume.

**Immunostaining**

Human donor eyes from wet AMD patients or age-matched controls were obtained from the National Disease Research Interchange (NDRI; Philadelphia, PA) with full ethical approval for use in research. NDRI obtains informed consent for all donor material (http://ndrisource.org/for-donors-patients) and all procedures conform to the principles set out in the WMA Declaration of Helsinki and the Department of Health and Human Services Belmont Report. All sample tissues were anonymized prior to receipt in the laboratory. Mouse eyes were harvested 14 days after L-CNV induction. The eyes were enucleated and fixed in 4% paraformaldehyde/PBS overnight. The anterior segment, lens, and vitreous humor were removed, and the posterior eye cups were prepared for standard paraffin sections or retinal flat mounts. Deparaffinized sections were treated with rodent deblocker (Biocare Medical, Concord, CA, USA) for antigen retrieval. The sections or flat mounts were washed with PBS then permeabilized with 0.3% Triton X-100 and nonspecific binding blocked by 10% normal goat serum plus 1% BSA in PBS. They then received primary antibody (polyclonal anti-FECH (clone C20)) at 1:500 for 16 h at 4°C. After primary incubation, sections or whole mounts were washed and incubated for 1.5 h at room temperature with secondary antibody (Cy3-conjugated goat anti-rabbit IgG, 1:600) at 4°C with 0.1% Triton X-100. We used a vascular-specific lectin (*Ricinus communis* agglutinin I; Vector Laboratories, Inc.) conjugated to Alexa Fluor 488 to label the retinal vasculature. This was incubated for 30 min at room temperature in 1:400 of 10 mM HEPES plus 150 mM NaCl and 0.1% Tween-20. After washing, specimens were mounted in aqueous mounting medium (VectaShield; Vector Laboratories, Inc.) and coverslipped for observation by confocal microscopy and quantification where noted by a masked investigator using ImageJ software. All microscopic images in a given panel were acquired with identical exposure settings.

**Choroidal sprouting assay**

Choroidal sprouting was assessed as described (Sulaiman et al., 2015). Briefly, pieces of choroid–sclera dissected from mouse eyes were embedded in Matrigel and grown in EGM-2 medium plus antibiotics for 72 h to allow sprouting to initiate. The indicated concentrations of griseofulvin (in DMSO, final DMSO concentration 1%) were added and growth allowed to proceed for 48 h. Images were taken and growth quantified by measuring the distance from the edge of the choroidal piece to the growth front in four directions per sample.

**Retinal endothelial cell sprouting assay**

An *ex vivo* murine retinal angiogenesis assay (EMRA) was performed according to the previously published protocol (Rezzola et al., 2013) with slight modifications. Briefly, retinal fragments were isolated from adult C57BL/6J mice (3.5–4.5 weeks old) and embedded in fibrin gel [containing bovine fibrinogen (4 mg/ml), aprotinin (5 µg/ml), and bovine thrombin (500 mU/ml)] in serum-free DMEM.
and allowed to gel at 37°C. After clotting for 20–30 min, 400 µl of DMEM containing 10% FBS and human VEGF165 (100 ng/well) was added and replaced after 3 days. After day 7, medium was replaced with DMEM containing 2% FBS plus VEGF165 (100 ng/well) and treated with griseofulvin (20 and 40 µM) or vehicle control (DMSO, 0.5% final concentration). Endothelial sprouting was monitored and photographed daily until day 3 after treatment. Image analysis of the retinal endothelial sprouting was performed manually using ImageJ software, and quantification was performed of least six regions of interest per treatment per time point. Endothelial cell sprouting was represented as sprouting distance in pixels.

**Griseofulvin feeding**

Mice were fed griseofulvin for a total of 3 weeks, with chow changed every 2–3 days. Standard mouse chow (5 g/mouse/day) was mixed in water (2.2 ml H2O/gram chow), soaked for 15 min, and then mashed. Griseofulvin doses were prepared at 0.0% (control), 0.5%, and 1.0% with 0.0, 0.5, and 1.0 g griseofulvin : 10 ml corn oil : 100 g mouse chow ratio, respectively. Both 0.5 and 1.0% doses were expected to substantially inhibit FECH and induce a protoporphyrin-like phenotype, based on previously published work (Lochhead et al, 1967; Holley et al, 1990; Martinez et al, 2009). During treatment, the mice were examined and weighed 3 times/week (Fig EV5A). On day 8, mice underwent L-CNV as above and were imaged by OCT at day 15 and day 22, at which time they were euthanized. The eyes were enucleated and flat mounts prepared as described above. The livers were dissected out and weighed (Fig EV5B).

**NOS assay**

NOS activity was measured in cells as described before (Räthel et al, 2003). Briefly, HRECs (10,000 cells/well) were seeded in 96-well clear-bottom black plates and incubated for 24 h at 37°C and 5% CO2. Cells were washed with PBS and incubated for 5 min at 37°C with 100 µl of 100 µM L-arginine prepared in PBS. Subsequently, 100 nM of DAF-2 diacetate was added to each well and fluorescence readings were taken at excitation and emission wavelengths of 495 and 515 nm, respectively, using the Synergy plate reader (Biotek).

**Hemin pull-down**

HRECs were grown in EGM-2 medium in 10-cm plates until they reached ~50% confluency. Cells were treated with 10 µM NMPP or DMSO control for 1 week. Medium was changed every 2 days with treatments added to the fresh medium every time. Cells were lysed with NP-40 lysis buffer (20 mM Tris, pH 8.0, 150 mM NaCl, 1% NP-40, 20 µM leupeptin, 1 mM PMSF, 1 mM NaF, 1 mM β-glycerophosphate, 2 mM sodium orthovanadate, 2 mM EDTA, 10% glycerol) and then centrifuged at 14,000 g for 10 min at 4°C. Supernatant was collected and samples were pre-cleared by incubation with Neutravidin beads for 1 h at 4°C followed by centrifugation at 500 g for 5 min at 4°C. Supernatant was collected, and protein concentration was determined using a Bradford assay. Equal amounts of total protein (40 µg) from each sample were incubated with ~50 µl hemin agarose beads, pre-washed three times with lysis buffer, for 1 h at 4°C.

The beads were then washed two times in NP-40 buffer, followed by two washes in high-salt buffer containing 20 mM Tris–HCl pH 8.0, 350 mM NaCl, and 1 mM PMSF. The beads were then washed once with a very high-salt buffer containing 20 mM Tris–HCl and 1 mM PMSF. The beads were then washed again in salt-free buffer containing 20 mM Tris–HCl and 1 mM PMSF. After 5-min incubation, the beads were collected and any residual liquid was removed using a Hamilton syringe. The hemin agarose beads were heated in 30 µl of 2× SDS–PAGE gel loading dye containing 30 µl of 2-mercaptoethanol for 10 min at 70°C to release the bound proteins. After heating, the contents were allowed to cool, and after a quick spin, the eluate was collected using a Hamilton syringe. Eluates were separated in a 4–20% gradient SDS–PAGE and then transferred onto PVDF membranes. Proteins were immunoblotted with antibodies against α-tubulin (1:1,000) or eNOS (1:500). All dilutions were made in Tris-buffered saline–0.05% Tween-20 buffer containing 2% bovine serum albumin (BSA).

**PPIX buildup assay**

HRECs were grown in a 6-well plate until confluent. Cells were serum starved overnight in EBM-2 medium. Fresh EGM-2 medium containing DMSO or cromistastranone (1) was added to cells, and they were incubated at 37°C for 1 h followed by the addition of 1 mM 5-ALA to increase flux through the heme biosynthetic pathway. After 3 h of incubation in the dark at 37°C, the cells were trypsinized and lysed in buffer containing 25 mM HEPES–NaOH pH 7.4, 150 mM NaCl, 1% NP-40, 10% glycerol, and 1 mM PMSF. The cell lysates were incubated in the dark at 4°C for 20 min on a shaker and centrifuged at 12,000 g for 15 min. Supernatants were collected for analysis. In a 384-well black plate, 20 µl of supernatant was mixed with 20 µl of 1:1 solution of 2 M perchloric acid and methanol. After 5 min of incubation, fluorescence readings were taken at excitation and emission wavelengths of 407 and 610 nm using the Synergy plate reader.

**Iron chelation**

Compounds (cromistastranone, EDTA, or deferoxamine) or DMSO (1 µl) were incubated with 2.5 mM freshly prepared ferrous ammonium sulfate in a final volume of 100 µl for 5 min at 37°C. Ferrozine (100 µl of 2.5 mM solution) was added to the wells, and spectrophotometric readings were taken at 562 nm using the Synergy plate reader. Decrease in absorbance readings at 562 nm represented the degree of iron chelation.

**Apoptosis assays**

The caspase-3 immunofluorescence assay was performed as described previously (Basavarajappa et al, 2014). Briefly, cells were plated on coated coverslips and incubated in EGM-2 medium overnight before treating with siRNAs. After 24 h of transfection, the cells were fixed in 4% paraformaldehyde and permeabilized using 0.5% Triton X-100 solution prepared in PBS. The cells were incubated with cleaved caspase-3 antibody (1:200 dilution) overnight at 4°C. Dylight 488-conjugated goat anti-rabbit secondary antibody (1:400) was used to probe the cleaved caspase-3 antibody. The
coverslips were mounted using Vectashield mounting medium containing DAPI for nuclear staining. The cells were imaged using an LSM 700 confocal microscope.

The TUNEL assay was carried out as described previously (Basavarajappa et al., 2015). Briefly, cells (25,000 per coverslip) were seeded on coverslips and 24 h later the cells were transfected with siRNAs or treated with the indicated compound concentrations for 24 h. Cells were then fixed in 4% paraformaldehyde for 20 min and permeabilized using 0.25% Triton X-100 prepared in PBS. Then, apoptotic cells were visualized using the Click-it TUNEL assay kit as per the manufacturer’s instructions, with DAPI counter-stain. The percentage of apoptotic cells was counted on three low-power fields per coverslip using ImageJ software.

qRT–PCR

The assay was performed as described previously (Basavarajappa et al., 2014). RNA was extracted from cells treated as indicated using Trizol. cDNA was synthesized from 500 ng RNA using random primers and iScript reverse transcriptase. qPCR was performed in 10 µl volumes in a 384-well plate, with Fast Advanced Master Mix and TaqMan probes on a ViiA7 thermal cycler (Life Technologies). Primer/probesets used were as follows: FECH (Hs01555261_m1), HIF1A (Hs00153153_m1), NOS3 (Hs01574659_m1), VEGFA (Hs00900055_m1), VEGFR2 (Hs00911700_m1), and housekeeping controls GAPDH (Hs99999905_m1), HPRT (Hs02800695_m1), and TBP (Hs00427620_m1). The data were analyzed using the ΔΔCt method (Livak & Schmittgen, 2001). The expression levels of genes were normalized to the 3 housekeeping genes and calibrated to the negative siRNA-treated sample.

Statistical analyses

Both in vitro and in vivo data were analyzed using unpaired Student’s t-test or one-way ANOVA with Dunnett’s or Tukey’s post hoc tests for comparisons between appropriate groups using GraphPad Prism v. 7.0 software. The choroidal and retinal sprouting assays were analyzed using two-way repeated measures ANOVA with Dunnett’s post hoc tests, while the knockdown time course was analyzed by two-way repeated measures ANOVA with Bonferroni’s post hoc tests. Two-sided P-values < 0.05 were considered significant in all tests.

Expanded View for this article is available online.

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Author contributions

TWC and HDB conceived the study and drafted the manuscript. HDB performed the in vitro experiments with the help of RSS, TS, KLS, CMB, BT, and MS. RSS and XQ performed the in vivo experiments with the help of JQ, SA, and KG. RSS and SSPB performed the sprouting assays. S-YS and XQ performed the sprouting assays. S-YS and BL synthesized the affinity reagents. MBG and MEB analyzed data and provided conceptual input. All authors discussed the results and commented on the manuscript.

Conflict of interest

H.D.B. and T.W.C. are named inventors on a patent application related to this work. The other authors declare no competing financial interests.

For more information

The Coron laboratory website: http://glick.iu.edu/corson
The FECH gene: http://www.ncbi.nlm.nih.gov/gene/2235
BrightFocus Foundation information on AMD: http://www.brightfocus.org/macular

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