Von Hippel-Lindau mutations disrupt vascular patterning and maturation via Notch

Alexandra Arreola,1 Laura Beth Payne,2 Morgan H. Julian,2,3 Aguirre A. de Cubas,4 Anthony B. Daniels,5,6,7 Sarah Taylor,2 Huaning Zhao,2,9 Jordan Darden,2,10 Victoria L. Bautch,1,11,12 W. Kimryn Rathmell,4,6,8 and John C. Chappell2,3,9,10

1Lineberger Comprehensive Cancer Center, University of North Carolina at Chapel Hill (UNC-CH), Chapel Hill, North Carolina, USA. 2Center for Heart and Regenerative Medicine and 3Department of Basic Science Education, Virginia Tech Carilion School of Medicine and Research Institute, Roanoke, Virginia, USA. 4Department of Medicine, Division of Hematology and Oncology, 5Department of Ophthalmology and Visual Sciences, 6Department of Radiation Oncology, and 7Department of Biochemistry, 8Department of Medicine, 9Department of Biomedical Engineering and Mechanics, 10Graduate Program in Translational Biology, Medicine, and Health, Virginia Polytechnic Institute and State University, Blacksburg, Virginia, USA. 11Department of Biology and 12McAllister Heart Institute, UNC-CH, Chapel Hill, North Carolina, USA.

Introduction
Von Hippel-Lindau (VHL) syndrome is an autosomal dominant inherited predisposition to the onset of cancer in neurological tissues, as well as in the kidney, adrenal gland, pancreas, and liver (1). Genetic lesions that compromise the normal VHL protein (pVHL) function can give rise to different types of carcinomas, such as clear cell renal cell carcinoma (ccRCC) (2). In addition to elevated risk for ccRCC, 60%–80% of patients with VHL syndrome, including those with type 2B missense VHL mutations, will develop hemangioblastomas (3–5). These benign neoplasms are composed of proliferating endothelial and stromal cells that arise in the brain and brain stem, as well as the spinal cord and retina, and often result in catastrophic effects for patients due to their site of presentation (6). Among other functions, the ubiquitin E3 ligase substrate recognition activity of pVHL provides essential modulation of hypoxia-inducible factors (HIFs), which orchestrate transcription of downstream gene targets in proportion to tissue oxygen levels. The VHL type 2B mutation R167Q has been described as a loss of pVHL interaction with Elongin C but
retention of partial regulation of HIF-1α and HIF-2α levels, which may help explain the observed clinical spectrum (7). Numerous HIF-1α targets are key regulators of blood vessel growth and remodeling, the most potent of which is VEGF-A (8).

Endothelial cells respond to VEGF-A through the receptor tyrosine kinase Flk-1/VEGF Receptor-2 (VEGFR-2), which promotes endothelial cell proliferation and sprouting migration (9). The Notch pathway intersects with VEGF-A signaling to coordinate endothelial cell behaviors such that endothelial “tip” cells emigrate outward from parent blood vessels, while “stalk” cells largely divide to promote vessel elongation (10, 11). Genetic loss of delta-like 4 (Dll4) or pharmacological blockade of Notch1 receptor signaling, such as through N-[N-(3,5-Difluorophenacetyl)-L-alanyl]-S-phenylglycine t-butyl ester (DAPT) treatment, leads to a hypersprouting phenotype and the formation of excessive vessel branches (10, 12), though this nascent vasculature is not always functional (13, 14). Endothelial cells also express the Notch ligand Jagged1 (Jag1), an antagonist of Dll4 that competes for binding of Notch1 and thus modulates tip cell formation and vascular network complexity (15). Notch signaling also specifies arterial-venous identity of endothelial cells (16), providing key signals for vessel maturation and mural cell development (17–19). For instance, endothelial Jag1 induces murine cell expression of vascular smooth muscle cell (SMC) proteins such as α-smooth muscle actin (αSMA) via binding of the Notch3 receptor (20, 21). These, and numerous other studies, therefore support a model in which elevated Notch signaling restricts overall branching complexity of a developing network while promoting vessel maturation and arterialization (22); thus, Notch blockade can increase vessel branching and limit processes involved in maturation, such as the acquisition of functional vascular SMCs (19, 21).

The effect of Notch blockade in the setting of variable levels of pathogenic pathway activation, however, is unknown.

Aberrant Notch signaling has been implicated in pathological conditions associated with VHL syndrome, most notably in the onset and progression of ccRCC (23, 24). While VHL mutations are known to disrupt the HIF/VEGF-A signaling axis and lead to abnormal vascular remodeling (25), the intersection with the Notch pathway, particularly in the vascular compartment, remains to be fully elucidated in the VHL mutation background. Recent studies suggest solid tumors frequently acquire a resistance to antiangiogenic, and specifically anti-VEGF, therapies (26, 27); therefore, a more complete understanding of Notch signaling in the setting of VHL loss or mutation is critical for developing alternative therapeutic targets within the Notch pathway for treating ccRCC and VHL-related conditions such as hemangioblastoma.

In the current study, we explored blood vessel development in the context of conditional biallelic \( Vhl \) loss (i.e., \( UBC^{CreER^{+/+}}/Vhl^{+/+} \)), as well as in the setting of a specialized model of type 2B \( Vhl \)-knock-in mutation, rendered monoallelic by conditional deletion of a WT copy of \( Vhl \) (i.e., \( UBC^{CreER^{+/+}}/Vhl^{fl/2B} \)), mimicking the loss of heterozygosity that takes place in human VHL disease (referred to as VHL type 2B mutant). Embryonic stem cell–derived (ES cell–derived) blood vessels harboring the type 2B G518A missense mutation in \( Vhl \) (equivalent to the arginine 167 to alanine \([R167Q]\) in humans) displayed a more severe vascular dysmorphogenesis than the \( Vhl^{+/−} \) vasculature. Endothelial cells isolated from both genotypic backgrounds, however, exhibited transcriptional alterations in VEGF-A and Notch pathway genes. In vivo analysis of the early stages of mouse retinal vessel development P5 revealed branching morphology defects in the setting of a \( Vhl \) type 2B mutation (i.e., \( UBC^{CreER^{+/+}}/Vhl^{fl/2B} \) and \( UBC^{CreER^{+/+}}/Vhl^{R167Q} \)) but only subtle changes in vessels of conditionally deleted \( Vhl \)-null mice (i.e., \( UBC^{CreER^{+/+}}/Vhl^{fl/0} \)). At slightly later stages of retinal development (P7), we found that genetic loss of \( Vhl \) (i.e., \( UBC^{CreER^{+/+}}/Vhl^{fl/0} \)) caused an aberrant increase in arterial maturation (e.g., premature elevation of pericyte αSMA expression) at the expense of expanding capillary density. \( Vhl \) type 2B mutation carriers induced to lose the WT copy of \( Vhl \) (i.e., \( UBC^{CreER^{+/+}}/Vhl^{fl/0} \)), however, exhibited this accelerated arterial formation without a concomitant decrease in retinal microvessels. Pharmacological inhibition of Notch signaling rescued the aberrant arterial maturation phenotype while causing significant increases in vessel branching. Applying RNA Sequencing (RNA-Seq) analysis to these \( VHL \) mutants, we found distinct changes in gene expression patterns of key signaling networks, including the Notch, HIF, SMC contraction, and FoxO/TGFβ pathways. Toward the completion of retinal development at P21, conditionally \( Vhl \)-null and type 2B mutant (i.e., \( UBC^{CreER^{+/+}}/Vhl^{fl/0} \) and \( UBC^{CreER^{+/+}}/Vhl^{fl/2B} \), respectively) vasculature displayed severe morphological defects, reflecting misregulation within the Notch pathway that integrates with and compounds defects in these other critical signaling cascades. Taken together, these results demonstrate that the type 2B \( VHL \) mutation may influence vascular changes and hemangioblastoma formation by accelerating the maturation of larger-caliber arteries in addition to maintaining and expanding capillary density, which could, in turn, worsen pathological progression via blood perfusion defects.
Results

Type 2B Vhl mutation disrupts VEGF and Notch signaling and causes blood vessel-branching defects. The VHL complex provides essential regulation of HIFs, which in turn modulate expression of a number of downstream proangiogenic target genes, including VEGF-A. Because Notch signaling intersects with the VEGF-A pathway (10, 11), we hypothesized that complete genetic loss of Vhl (Vhl–/–) and the type 2B Vhl mutation cause vessel overgrowth through downstream disruptions in both the VEGF-A and Notch pathways. Endothelial cells that arise during mouse ES cell differentiation undergo angiogenic sprouting and form lumensed blood vessels, akin to vascular development in vivo (28, 29). We analyzed WT ES cell–derived vascular networks for their overall branching complexity and compared them with vessels formed in the Vhl-null (Vhl–/–) and –type 2B mutant backgrounds (Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.92193DS1). Interestingly, Vhl–/– vessels did not exhibit any significant changes in overall vessel branching. In contrast, type 2B mutant (i.e., Vh2B/2B) vasculature showed a significant increase in branching complexity, suggesting that the type 2B Vhl mutation has distinct effects on the molecular mechanisms governing blood vessel branching.

To further test our hypothesis regarding VEGF-A and Notch pathway disruptions downstream of genetic Vhl mutations, we analyzed the transcriptional profile of WT, Vhl–/–, and Vh2B/2B endothelial cells by assessing candidate targets within these pathways. We enzymatically dissociated ES cell cultures from each genetic background and used magnetic bead–assisted cell sorting (i.e., via platelet-endothelial cell adhesion molecule-1 [PECAM-1]/CD31 labeling) to enrich for endothelial cells. We assessed mRNA levels by quantitative PCR (qPCR) specifically evaluating VEGF-A and VEGFR2/Flk-1 levels, as well as the Notch pathway components Dll4, Jag1, Notch3, and Hey2 (Supplemental Figure 1). Although VEGFR2/Flk-1 mRNA levels were significantly higher in both Vhl null mutant backgrounds, only the type 2B Vhl mutant endothelial cells harbored a significant increase in VEGF-A transcript levels. Interestingly, Dll4 mRNA levels were significantly lower in Vhl–/– endothelial cells, as compared with WT, while these transcripts were significantly elevated in the type 2B mutants. Endothelial Notch3 expression was significantly higher in both backgrounds, and Jag1 expression was raised in both but showed a significant increase only in the Vhl–/– context. The Notch pathway transcription factor Hey2 was significantly elevated in endothelial cells only expressing type 2B Vhl (i.e., Vh2B/2B), however, it was unchanged with full genetic loss of Vhl. Taken together, these observations suggest that the Vhl–/– and type 2B Vhl mutations elicit specific effects on endothelial cells, which is reflected in expression of VEGF-A and Notch pathway components, and these distinct signaling environments yield vascular networks with overall unique morphologies.

Conditional Vhl mutations induce abnormal blood vessel development in early-stage postnatal mouse retinas. In observing differential effects of Vhl–/– and Vh2B/2B on vascular development, we hypothesized that in vivo blood vessel formation would also exhibit distinct abnormalities due to disruption of the VEGF-A and Notch signaling pathways. To test this hypothesis, we assessed blood vessel formation in the developing retinas of P5 mice harboring (i) tamoxifen-inducible Cre-recombinase activity (i.e., UBCCreER+/+) and (ii) loxp sites for creating mutated Vhl alleles via targeted recombination (Vhlfl/fl and Vh2B/2B; Supplemental Figure 2). After tamoxifen administration from P1–P3, retinas were collected at P5 and immunostained to visualize the developing vascular networks using confocal microscopy. Arterial and venous vasculature (in the Vhl type 2B mutant retinas i.e., UBCCreER+/+Vhl2B/2B and UBCCreER+/+Vhl2B/2B) had significant elevations in branching complexity compared with littermate controls (i.e., UBCCreER+/+Vhfl/fl) (Figure 1). Interestingly, retinal vessels in Vhl heterozygous (UBCCreER+/+Vhl2B/2B) mice exhibited a significant increase in arterial, but not venous, branching. Consistent with the Vhl–/– ES cell–derived vessels, the induced that Vhl-null retinas (i.e., UBCCreER+/+Vhfl/fl) lacked vessel-branching changes. Furthermore, conditional type 2B mutant vessels (Vhl type 2B mutant; i.e., UBCCreER+/+Vhl2B/2B) were more limited in their expansion across the retinal surface, as were the UBCCreER+/+Vhl2B/2B and UBCCreER+/+Vhl2B/2B vasculature, though to a lesser extent. Arterial sprouting in littermate control retinas (i.e., UBCCreER+/+Vhfl/fl) was significantly (1.5-fold) higher than sprout formation within venous regions; however, this difference was lost with all induced Vhl mutations (UBCCreER+/+Vhl2B/2B and UBCCreER+/+Vhl2B/2B), suggesting a potential shift from predominant angiogenic sprouting and moderate vessel maturation to a balance between these 2 stages of vessel formation. Overall, these data are consistent with the idea that the branching complexity of developing vessels is severely affected by the type 2B Vhl mutation, such that even in the presence of the WT Vhl gene, we observe disrupted vascular development (Vhfl/fl heterozygotes), while heterozygous or homozygous deletions of the Vhl gene are less disruptive to normal blood vessel formation.
Notch inhibition rescues maturation defects in the retinal vasculature of postnatal Vhl mutant mice. Vessel-branching irregularities observed in the development of conditional Vhl mutant retinas suggested that additional aspects of arterial and venous network formation might also be perturbed. Because Notch signaling is known to be involved in arterial-venous specification and vessel maturation (16–18, 20), we hypothesized that later stages of vascular development might also be adversely affected by the induction of genetic Vhl mutations. Therefore, we analyzed the developing retinal vasculature of P7 conditional Vhl mutant mice for branching complexity and hallmarks of arterial maturation, specifically the morphology and spatial distribution of αSMA+ vessels. In addition, we explored the vessel maturation role of Notch signaling downstream of Vhl mutations by pharmacologically blocking the Notch pathway in each of the Vhl mutant backgrounds (P5–P6 i.p. administration of the γ-secretase inhibitor DAPT; Supplemental Figure 2), aiming to observe a potential rescue in branching and arterial maturation defects. Vessel branching within arterial and venous regions was significantly reduced in conditional Vhl-null retinas (UBCreER+/+Vhlf/fl; Figure 2), comparable with the sparsely branched retinal vasculature of mice with a gain-of-function mutation for Notch signaling (i.e., conditional overexpression of Notch1 intracellular domain [N1ICD], UBCcreER+/+ RosaN1ICD; Supplemental Figure 3). In contrast, induction of the type 2B Vhl mutation (UBCreER+/+Vhlf/2B) had little to no effect on branching (Figure 2). Notch inhibition elevated retinal vessel branching within all experimental groups, regardless of association with arterial and venous areas. In spite of these increases, UBCcreER+/+Vhlf/fl vasculature was not fully rescued back to littermate control branching levels by blocking Notch signaling. These observations...
again underscore that the type 2B \(Vhl\) mutation uniquely affects blood vessel formation, consistent with the previously described data from the ES cell–derived vessels and the P5 retinal vasculature.

To compare the influence of the different \(Vhl\) mutations on vessel maturation, we assessed P7 retinal vasculature for the morphology and spatial distribution of \(\alpha\text{SMA}^+\) vessels. Specifically, we measured (i) the percent length of primary arterial vessels covered with cells expressing \(\alpha\text{SMA}\), (ii) the diameter of these primary retinal arteries/arterioles, and (iii) the relative amount of \(\alpha\text{SMA}\) expression aberrantly occurring beyond the primary retinal arteries (i.e., within the capillary plexus region at the vascular front) (Figure 3). In contrast to vessel-branching changes, arterial morphologies were perturbed in both induced \(Vhl\)-null and –type 2B mutations (\(UBC^{CreER+}/Vhl^{fl/fl}\) and \(UBC^{CreER}/Vhl^{fl/2B}\), respectively). Primary arteries within these \(Vhl\) mutant retinas displayed \(\alpha\text{SMA}\) expression extending further toward the vascular front, and these arteries had significantly larger diameters compared with the littermate control retinas (both \(UBC^{+/+}/Vhl^{fl/fl}\) and \(UBC^{+/+}/Vhl^{fl/2B}\)). Strikingly, \(\alpha\text{SMA}\) expression along microvessels at the periphery of the \(Vhl\)-null and –type 2B mutant retinal vasculature exhibited an abnormal spatial distribution. In particular, more perivascular cells at the vascular front expressed a higher level of \(\alpha\text{SMA}\) than is typically displayed at this developmental time point (21), and this excess of \(\alpha\text{SMA}\) was not found in nonmutant littermate controls harboring at least
1 WT Vhl allele (i.e., UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/fl} and UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/2B}) (Figure 3). Notch inhibition rescued all of the defects observed in the morphology and spatial distribution of αSMA\(^+\) vessels of P7 retinal vasculature. Taken together, these results, which are summarized in Supplemental Figure 4, suggest that Notch signaling makes critical contributions to developmental maturation of arterial vessels in the setting of Vhl deficiency.

Vascular pericytes have been suggested to regulate angiogenic remodeling at the retinal vascular front, specifically by influencing endothelial cell interactions via Notch signaling (30). In addition, recent studies have suggested that coronary SMCs can arise during development from pericytes expressing neural glial antigen 2 (NG2) (31). Thus, we asked if the perivascular cells with increased αSMA expression in

Figure 3. In P7 mouse retina, conditional Vhl-null and –type 2B mutations accelerate vessel maturation via excess alpha-smooth muscle actin expression, and this defect can be rescued by blocking Notch signaling. (A–H) Representative images of P7 mouse retinal vasculature stained with isolectin B4 (top rows) and αSMA (middle rows). A quarter region, or leaflet, is shown for UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/fl}, UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/2B}, UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/fl}, and UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/2B} littermate retinas exposed to tamoxifen and subjected to either control treatment (DMSO, A–D) or Notch inhibition (DAPT, E–H). Scale bars: 200 μm. (I) Percent length of primary artery covered by αSMA\(^+\) cells in control (triangles) and Notch-inhibited (squares) P7 vascular networks. *P ≤ 0.05 vs. control UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/fl} and UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/2B}, and vs. Notch-inhibited within same genotype. (J) Diameter of primary retinal artery measured from control (triangles) and Notch-inhibited (squares) P7 vascular networks. *P ≤ 0.05 vs. control UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/fl} and UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/2B}, and vs. Notch-inhibited within same genotype. (K) Percent mislocalization of αSMA expression not associated with primary retinal arteries in control (triangles) and Notch-inhibited (squares) retinas. *P ≤ 0.05 vs. control UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/fl} and UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/2B}, and vs. Notch-inhibited within same genotype. **P ≤ 0.05 vs. control UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/fl}, and vs. Notch-inhibited UBC\textsuperscript{CreER}/-- Vhl\textsuperscript{fl/2B}. Values are averages ± SEM. n = 6–8 retina images per group (i.e., 3–4 from 2 experimental litters). All statistical comparisons performed using 1-way ANOVA followed by pair-wise comparisons with a 2-tailed Student t test.
the conditional Vhl mutant retinas were also positive for NG2, a well-established retinal pericyte marker (32). Indeed, we found a substantial overlap in the αSMA and NG2 signals (Figure 4), suggesting that the genetic loss of the Vhl gene (UBC\textsuperscript{CreER/+}Vhl\textsuperscript{fl/fl}) and, to a greater extent, the type 2B Vhl mutation (UBC\textsuperscript{CreER/+}Vhl\textsuperscript{fl/2B}) accelerates vascular pericyte expression of αSMA that likely yields an increased population of contractile vascular SMCs.

Notch signaling is among several vascular morphogenesis pathways disrupted by inducing Vhl loss or type 2B mutation. The pVHL regulates HIF activity, which in turn modulates numerous downstream vascular morphogenesis cues including VEGF-A and PDGF, among several others (8). While our data support Notch signaling as an additional downstream effector of pVHL activity, we hypothesized that the Vhl-null and –type 2B mutation scenarios (i.e., UBC\textsuperscript{CreER/+}Vhl\textsuperscript{fl/fl} and UBC\textsuperscript{CreER/+}Vhl\textsuperscript{fl/2B}, respectively) likely elicit critical changes in gene expression within other vascular morphogenesis pathways, in addition to Notch. To test this idea, we performed transcriptome analysis of whole retinas from P7 conditional Vhl-null and –type 2B mutation mice (and control littermates). RNA-Seq data (uploaded to NCBI Gene Expression Omnibus; GEO) was mapped to the mouse reference genome, and we conducted pathway analysis using the Database for Annotation, Visualization and Integrated Discovery (DAVID) (33, 34). In addition to significant changes in the expression of Hey2, an important transcriptional target in the Notch pathway (Figure 5), we found disrupted transcription of components within other pathways as well. Among several effectors altered within the HIF pathway, Egl-9 Family Hypoxia Inducible Factor 3 (Egln3) was upregulated in the null (UBC\textsuperscript{CreER/+}Vhl\textsuperscript{fl/fl}) and type 2B (UBC\textsuperscript{CreER/+}Vhl\textsuperscript{fl/2B}) retinas, consistent with VHL/HIF signaling dynamics in hypoxia (35). Also significantly

Figure 4. Pericytes at the vascular front of P7 conditional Vhl-null and Vhl type 2B mutant retina vessel networks ectopically express αSMA but not in littermate control vessels. (A–F, H–M, O–T) Representative images of P7 mouse retinal vasculature stained for isolectin B4 (A, H, O), αSMA (B, I, P), and neural glial antigen-2 (NG2) (C, J, Q). The NG2 image is merged with the isolectin B4 image (D, K, R), with the αSMA image (E, L, S), and with both isolectin B4 and αSMA (F, M, T). Arrows indicate NG2+ cells overlapping with αSMA signal. The dotted lines in E, L, and S indicate the region of interest where pixel intensity values were measured and normalized. Scale bars: 50 μm. (G, N, U) Pixel intensity values for each line scan of NG2 and αSMA signals in the region indicated by the dotted line drawn in E, L, and S, respectively.
Elevated was the expression of genes involved in vascular smooth muscle contraction, such as rho guanine nucleotide exchange factor 12 (Arhgef12), which contributes to myosin regulation in SMCs (36). Transcriptional regulation within the FoxO/TGFβ signaling axis was also severely affected, as seen in reduced expression of Gadd45b. This protein induces growth arrest and can limit downstream TGFβ signaling (37); thus, reduced Gadd45b expression in the Vhl-null and –type 2B backgrounds suggests a lack of cell growth arrest and a permissive environment for TGFβ signaling, which is known to promote SMC differentiation (38, 39).

These transcriptional changes, alongside our observation of accelerated arterialization of the retinal vasculature in Vhl-null and –type 2B conditional mutants (UBC CreER/+ Vhlfl/fl and UBCCreER/+ Vhlfl/2B), lend further support to the idea that these Vhl mutations contribute to vascular defects encompassing misregulation of angiogenic remodeling, as well as of vessel maturation.

Retinal blood vessel defects in Vhl-null and –type 2B mutant mice worsen by P21. To determine if the vascular defects in the Vhl genetic loss and –type 2B mutation mice (UBC CreER/+ Vhlfl/fl and UBCCreER/+ Vhlfl/2B) persisted beyond early postnatal stages, we collected retinas from P21 mice (weaning) following the induction of these mutations from P1–P3. We found gross vascular dysmorphogenesis in the Vhl-null and –type 2B mutant retinas, including vessel patterning defects and extensive mis-localization of αSMA along retinal arteries and veins (Figure 6). Type 2B mutant vasculature (UBC CreER/+ Vhlfl/2B) appeared most severely affected, as these retinal vessels were more randomly distributed and exhibited aberrant αSMA expression along smaller diameter arterioles and arterial branches that connected directly to venules. While our conditional Vhl loss and –type 2B mutation models do not replicate
the full extent of the clinical situation, the vessel-branching and maturation phenotypes from the mouse data overlap with certain aspects of VHL-related retinopathy in patients (our unpublished data), warranting further investigation into how these vascular defects may give rise to clinical features of VHL-related retinopathy such as retinal capillary hemangiomas, exudative/tractional detachment, and scar tissue formation. Furthermore, they also support a model in which lesions in the VHL gene give rise to distinct vascular pathologies by disrupting vessel patterning mechanisms prior to the development of hemangioblastomas, such as those governed by the Notch pathway as well as those identified from our transcriptome analysis.

**Discussion**

VHL mutations cause defective blood vessel formation, as is often seen in the clinical presentation of hemangioblastomas within neural tissue of affected patients. In the present study, we show that conditional genetic loss of Vhl (UBC<sup>CreER<sup>+/+</sup>Vhl<sup>fl/fl</sup>) and the type 2B Vhl mutation (UBC<sup>CreER<sup>+/+</sup>Vhl<sup>fl/2B</sup>) have differential effects on vascular development via downstream perturbations in Notch pathway signaling and in conjunction with other pathway misregulation. Endothelial Notch signaling plays a critical role in sprouting angiogenesis (10–12), in vessel maturation through arterial-venous specification (16), and in the induction of contractile vascular SMC differentiation (17, 18, 20, 21). Our observations suggest Vhl lesions disrupt Notch signaling such that processes underlying vessel maturation are accelerated; most prominent among these...
is an altered spatiotemporal expression of αSMA. The early stages of vessel formation and branching are compromised during Vhl loss (UBCcreER+/Vhlfl/fl) but not with the type 2B Vhl mutation (UBCcreER+/Vhlfl/fl). These data suggest that Vhl mutations may not only cause substantial changes in angiogenic branching dynamics, but also strongly influence how blood vessels mature.

The pVHL provides essential regulation of HIFs, which in turn modulate downstream signals that promote vessel growth and remodeling such as VEGF-A. Kurihara et al. explored the VHL/HIF/VEGF signaling axis in postnatal retinas of mice with conditional Vhl loss (25). To induce cre/ERT2 expression in the retina, the authors used the Pax6 promoter, which is present in retina neural progenitor cells but may also be expressed by endothelial cells during early neural tissue development (40). Interestingly, they found that conditional Vhl-KO induced ectopic VEGF-A expression, but retinal blood vessel branching actually decreased, as was observed in the present study. Kurihara and colleagues found that diminished vascular branching resulted largely from vessel regression, which likely represents the vessel pruning actually decreased, as was observed in the present study. Kurihara and colleagues found that diminished vascular branching resulted largely from vessel regression, which likely represents the vessel pruning actually decreased, as was observed in the present study. Kurihara and colleagues found that diminished vascular branching resulted largely from vessel regression, which likely represents the vessel pruning actually decreased, as was observed in the present study.

Misregulation of the Notch signaling pathway has been implicated as a key factor in VHL syndrome manifestations and, most notably, in ccRCC onset and progression. Recent studies have shown an upregulation of the Notch1 receptor in human ccRCC, as well as increased levels of the cleaved intracellular region of Notch1 responsible for downstream signaling (23). Expression of the Notch ligand Jag1 is also highly upregulated in human ccRCC cells (24). While these changes in the Notch pathway have been implicated as contributing factors to renal cell transformation and ccRCC pathogenesis, disruptions in Notch signaling within vascular tissue may also influence disease progression by exacerbating pathological blood vessel remodeling. Our observations suggest that Vhl mutations likely influence vascular patterning in a way that undermines the Notch pathway, among several other effector pathways, such that blood vessel maturation is accelerated toward an arterial phenotype. With the formation of larger-caliber vessels, which presumably enhance blood flow to downstream tissues, it is intriguing to speculate that these changes may be an important factor in the aggressive growth and metastatic potential of ccRCC tumors (42). Thus, targeting the Notch pathway may rescue oncogenic defects, such as cell cycle misregulation (24), while also limiting tumor growth and metastasis by restricting the expansion of tumor-associated arterial vessels.

In the developing retina, vascular pericytes ensheath newly formed blood vessels to promote their stability and integrity. Pericytes can also give rise to arterial SMCs via Jag1/Notch3 signaling, as shown recently in the developing coronary arteries (31). The contractile capacity of capillary pericytes remains an open question (43, 44); however, in the current study, we found ectopic αSMA expression within NG2+ pericytes at the periphery of the vasculature within conditional Vhl-null and –type 2B mutants (i.e., UBCcreER+/Vhlfl/fl and UBCcreER+/Vhlfl/fl, respectively). Mural cells at the vascular front of littermate control retinas (i.e., UBCcreER+/Vhlfl/fl and UBCcreER+/Vhlfl/fl) lacked αSMA expression at this time point. This aberrant expression of αSMA in Vhl mutant animals suggests an accelerated maturation of retinal pericytes, presumably into precursors for SMCs. The differential effects of Vhl-null and –type 2B mutant Vhl retinal models are particularly interesting in light of the broad range of phenotype in patients with VHL syndrome with regard to retinal involvement and disease severity. Notch inhibition via DAPT treatment rescued this phenotypic abnormality, consistent with the notion that the Vhl mutations augment Notch pathway signals underlying SMC differentiation (17, 18, 20, 21). Notch signaling perturbations likely compound, and potentially intersect with, other cues downstream of pVHL activity, such as PDGF-BB (45) and TGFβ (39), both of which are known to promote SMC differentiation. Future work will be needed to extend the pathway analysis from the conditional Vhl-null and –type 2B mutant transcriptional profiles to elucidate the crosstalk between the Notch pathway and other signaling axes involved in the observed vascular dysmorphogenesis. Nevertheless, our results therefore implicate the Notch pathway as a potential therapeutic target for treating not only early-stage vascular malformations in patients with VHL syndrome, but also those defects affecting blood vessel maturation.
Deletion and type 2B VHL mutations underlie an increased risk for hemangioblastoma and ccRCC formation, owing in part to altered HIF expression (1) as well as defects in numerous other downstream pathways. While both VHL mutations cause abnormal, and often excessive, blood vessel remodeling in the clinical setting, our data suggest that these genetic lesions likely have differential effects on the nature of these vascular changes. Vhl-null mutation (UBC<sup>C</sup>/ERT2+/Vhl<sup>B</sup>/6) led mostly to accelerated vessel maturation, while the type 2B Vhl mutation (UBC<sup>C</sup>/ERT2+/Vhl<sup>B</sup>/28) caused an additional increase in vessel-branching complexity. The previously described partial regulation of HIFs by the mutant 2B Vhl (7) likely gives rise to a unique VHL/HIF downstream gene target axes affecting the aberrant vascular formation observed in type 2B mutants not commensurate in the Vhl-null setting. These results also may help explain the heterogeneity observed in the vascular architecture and integrity within ccRCC tumors (46), as well as the apparent discrepancy between levels of angiogenic factors, tumor vessel content, and tumor growth (28). This disconnect between proangiogenic signals and vessel outcomes has been linked to disruptions in the Notch pathway (14, 47). Our data also strongly implicate compromised Notch signaling downstream of Vhl mutations as a prominent molecular effector for the vascular dysmorphogenesis observed. Thus, developing therapeutic targets within the Notch pathway could offer novel treatment options for patients with VHL syndrome, especially those suffering from recurrent hemangioblastoma formation and ccRCC.

**Methods**

**Mutant mice and inducible genetic recombination experiments.** Mice expressing Cre-recombinase ubiquitously (Tg[UBC-cre/ERT2], The Jackson Laboratory, catalog 007001; denoted in the current study as UBC<sup>C</sup>/ERT2<sup>C</sup>/ERT2 mice; and ref. 48). UBC<sup>C</sup>/ERT2<sup>C</sup>/ERT2 mice generated previously (4) were bred with Vhl<sup>B</sup>/6 males (gift from Volker Haase, Vanderbilt University Medical Center) to yield Vhl<sup>B</sup>/6 offspring. Breeding UBC<sup>C</sup>/ERT2<sup>C</sup>/ERT2+Vhl<sup>B</sup>/6 mutants with Vhl<sup>B</sup>/28 females produced experimental litters with genotypes: (i) nonmutant littermate control, UBC<sup>C</sup>/ERT2<sup>C</sup>/ERT2/+Vhl<sup>B</sup>/6; (ii) littermate control with 1 WT Vhl allele and one type 2B mutant Vhl allele, UBC<sup>C</sup>/ERT2<sup>C</sup>/ERT2+/Vhl<sup>B</sup>/28; (iii) conditional Vhl-null, UBC<sup>C</sup>/ERT2<sup>C</sup>/ERT2+/Vhl<sup>B</sup>/6; and (iv) conditional Vhl type 2B mutant, UBC<sup>C</sup>/ERT2<sup>C</sup>/ERT2+/Vhl<sup>B</sup>/28. Where possible, animals were distributed across sex to avoid the introduction of sex bias. Pups were i.p. injected each day with 1 mg/ml of tamoxifen (Thermo Fisher Scientific) from P1–P3. Retinal harvest occurred on P5, P7, and P21 (Supplemental Figure 2). Cre-recombination activity following tamoxifen exposure was verified using a Cre reporter mouse line (Supplemental Figure 5). For Notch inhibition experiments, DAPT (Calbiochem) was dissolved in DMSO (MilliporeSigma), and 50 mg per kg of animal weight was administered i.p. on P5 and P6, with retina harvest occurring on P7 (Supplemental Figure 2). Equivalent volumes of DMSO-only were injected at the same time points for vehicle control animals.

**Retina immunostaining, imaging, and analysis.** Eyes were fixed by whole animal perfusion with 0.5% paraformaldehyde (PFA; Electron Microscopy Services) followed by immersion in 2% PFA for 2 hours at room temperature. Following PBS (VWR) rinse, retinas were dissected from the whole eye, immersed in 100% cold ethanol for 30 minutes, rinsed with PBS containing 1% Triton X-100 (PBS-T; Thermo Fisher Scientific) for 30 minutes at room temperature, and then blocked in 3% donkey serum (Jackson ImmunoResearch) in PBS-T for 1 hour at room temperature. Primary antibody incubation was performed in PBS-T overnight at 4°C with isoelectric GS-IB4 conjugated to AlexaFluor 488 (Molecular Probes, I21411, 1:100), αSMA conjugated to Cy3 (clone 1A4, MilliporeSigma, C6198, 1:200), and NG2 (MilliporeSigma, AB5320, 1:200). Secondary antibody incubation was also in PBS-T overnight at 4°C with donkey anti–rabbit AlexaFluor 647 (Jackson ImmunoResearch, 711-605-152, 1:400). Retinas were washed for 5 minutes 3 times in PBS-T, flattened, and mounted on slides in 50:50 PBS/glycerol (MP Biomedicals) with 1.5 coverslips. Images of retina whole mounts were acquired with a Zeiss LSM 880 confocal microscope with a ×40 objective, and 10–15 z-axis confocal scans were acquired and compressed. ImageJ was used to measure: (i) vessel-branch point densities, as described in the Supplemental Methods; (ii) the percentage of the retina radial to the optic disc occupied by vasculature (i.e., radial expansion); (iii) the number of vessel sprouts per length of the vascular front; (iv) the percent length of the primary arteries covered by αSMA+ cells; (v) the diameters of primary retinal arteries; (vi) the percent of αSMA+ cells not associated with the primary artery (i.e., mislocalized); and (vii) pixel intensity values across a line region of interest (i.e., line scan tool) (Figure 4).

**Retina transcriptome analysis.** Following induction of the genetic mutations as described above, retinas were collected from P7 mice and immediately digested in TRIzol (Invitrogen) to extract mRNA transcripts (n = 2 for each genotype). RNA was further isolated with 1-Bromo-3-chloropropane (MilliporeSigma) and...
ethanol and purified with an RNEasy Plus Kit (Qiagen). Using an Agilent Bioanalyzer, the Virginia Tech Biocomplexity Institute ensured the isolated mRNA was of a sufficiently high quality for RNA-Seq. Eight samples (4 genotypes × 2 replicates per genotype) were sent to GeneWiz for additional quality assessment, cDNA library generation, and sequencing on an Illumina HiSeq with a 2 × 150 configuration.

Sequencing reads were aligned against the Mouse GENCODE genome, Version M14 (January 2017 freeze, GRCm38, Ensembl 89) using the Spliced Transcripts Alignment to a Reference (STAR) software (49, 50). Reads were preprocessed and indexed using SAMtools (51). Normalization and differential expression was performed using DESeq2 (53). The top most significantly differentially expressed genes (FDR < 0.01 and log difference greater than 0.5 in magnitude) were considered for subsequent functional enrichment using DAVID bioinformatics tools (33, 34). The top 200 most differentially expressed genes were used for unsupervised hierarchical cluster analysis and visualized using heatmap representations. Raw data files are available upon request or from NCBI GEO (accession number GSE109102).

Statistics. Statistical analysis was conducted using GraphPad Prism 6. Multiple measurements were made for each parameter, and an average and SD were calculated from these values. Statistical comparisons were made using 1-Way ANOVA followed by pair-wise comparisons by 2-tailed Student t test. A P value less than 0.05 was considered significant.

Study approval. All animal experiments were conducted with review and approval from the UNC-CH and Virginia Tech IACUC. All protocols are reviewed and approved by IACUC boards. The UNC-CH NIH/PHS Animal Welfare Assurance Number is A3410-01 (expires 4/30/2021). The Virginia Tech NIH/PHS Animal Welfare Assurance Number is A-32081-01 (expires: 7/31/2021).

Author contributions
Research studies were designed by AA, LBP, VLB, WKR, and JCC. AA and JCC conducted experiments, and AA, ST, LBP, MHJ, and JCC acquired the primary data. Data analysis was performed by AA, AADC, HZ, JD, and JCC. AA, VLB, WKR, ABD, and JCC prepared the manuscript.

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Address correspondence to: John C. Chappell, Virginia Tech Carilion Research Institute, 2 Riverside Circle, Roanoke, Virginia 24016, USA. Phone: 540.526.2219; Email: JChappell@vtc.vt.edu. Or to: W. Kimryn Rathmell, Vanderbilt University, 798 Preston Research Building, 2220 Pierce Avenue, Nashville, Tennessee 37232-6307, USA. Phone: 615.936.3320; Email: Kimryn.Rathmell@Vanderbilt.edu.

AA’s present address is: Department of Pathology and Laboratory Medicine McLendon Clinical Laboratories, UNC Hospital, Chapel Hill, North Carolina, USA.


