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The adult kidney plays a central role in erythropoiesis and is the main source of erythropoietin (EPO), an oxygen-sensitive glycoprotein that is essential for red blood cell production. Decreases of renal pO2 promote hypoxia-inducible factor 2mediated (HIF-2-mediated) induction of EPO in peritubular interstitial fibroblast-like cells, which serve as the cellular site of EPO synthesis in the kidney. It is not clear whether HIF signaling in other renal cell types also contributes to the regulation of EPO production. Here, we used a genetic approach in mice to investigate the role of renal epithelial HIF in erythropoiesis. Specifically, we found that HIF activation in the proximal nephron via induced inactivation of the von Hippel–Lindau tumor suppressor, which targets the HIF-α subunit for proteasomal degradation, led to rapid development of hypoproliferative anemia that was associated with a reduction in the number of EPO-producing renal interstitial cells. Moreover, suppression of renal EPO production was associated with increased glucose uptake, enhanced glycolysis, reduced mitochondrial mass, diminished O₂ consumption, and elevated renal tissue pO₂. Our genetic analysis suggests that tubulointerstitial cellular crosstalk modulates renal EPO production under conditions of epithelial HIF activation in the kidney.





Renal epithelium regulates erythropoiesis via HIF-dependent suppression of erythropoietin

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Introduction

Renal oxygen sensing mechanisms play a key role in the regulation of erythropoiesis in adults, as the kidney is the main source of erythropoietin (EPO), an oxygen-sensitive glycoprotein hormone that is essential for normal erythropoiesis. EPO regulates rbc mass by promoting the survival of CFU-erythroid (CFU-E) cells and erythroblasts, which have not yet begun to synthesize hemoglobin (Hb) (1, 2). The induction of renal EPO synthesis represents a classic response to systemic hypoxia and results in increased rbc production, augmenting the oxygen-carrying capacity of blood when arterial pO₂ (PaO₂) is low. Failure to produce adequate amounts of EPO results in anemia, which is a common clinical feature of advanced chronic kidney disease (CKD) and is typically treated with recombinant EPO. Conversely, dysregulation of renal oxygen sensing, either by pharmacologic means or as a result of genetic mutations, can lead to inappropriately high EPO production and polycythemia (3).

The hypoxic induction of renal EPO is controlled by hypoxia-inducible factor 2 (HIF-2) (4). HIF transcription factors consist of an oxygen-sensitive α subunit (HIF-1 α , HIF-2 α [EPAS-1], and HIF-3 α) and a constitutively expressed β subunit, HIF- β , also known as the aryl-hydrocarbon receptor nuclear translocator (ARNT). HIF activity is controlled by oxygen-, iron-, and ascorbate-dependent prolyl-4-hydroxylase domain-containing proteins 1–3 (PHD1–3), which use 2-oxoglutarate (OG) as substrate for the hydroxylation of specific

proline residues within HIF- α . Under normoxia, HIF- α is hydroxylated and binds to the von Hippel–Lindau–E3 (VHL-E3) ubiquitin ligase complex, which subsequently leads to its proteasomal degradation (5). Under hypoxic conditions, however, HIF prolyl-4-hydroxylation is inhibited and HIF- α is no longer degraded and translocates to the nucleus, where heterodimerization with HIF- β /ARNT leads to transactivation of HIF target genes.

EPO synthesis in the kidney is dependent on tissue pO, (PtO2), and renal EPO production is controlled by the number of renal EPO-producing cells (REPCs), i.e., under hypoxic conditions, the number of REPCs increases proportionally to the degree of systemic hypoxia (6, 7). REPCs are contained within the peritubular interstitial/perivascular fibroblast population and express both pericyte markers (e.g., PDGFR-β polypeptide and ecto-5'-nucleotidase/CD73) and neuronal markers (e.g., microtubule-associated protein 2 and neurofilament protein light polypeptide) (7-9). They reside within the peritubular interstitial space of the deep cortex and outer medulla and are characterized by dendritic cellular projections, which form direct contacts with renal tubules and capillaries (7). Thus, intercellular crosstalk is likely to affect EPO synthesis in REPCs via paracrine and other signaling mechanisms or direct cell-cell interactions (10). However, these signaling mechanisms are not well understood. The degree of renal tubular workload has previously been suggested as modulating EPO production during hypoxia, presumably through effects on renal O, consumption. Serum EPO (sEPO) levels were diminished in rats treated with acetazolamide, but not with furosemide. Both are diuretics that decrease tubular sodium reabsorption and thus decrease renal O, consumption

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(11). A negative feedback loop also appears to operate during ascent to high altitude, where sEPO levels decrease before a significant increase in rbc mass is observed (12, 13). The molecular basis of this feedback regulation is not clear.

While it is well documented that hypoxia induces EPO via HIF-2 activation in REPCs, the role of hypoxia/HIF signaling in other renal cell types in the regulation of renal EPO synthesis is unclear. Here, we used a tetracycline-inducible genetic model of pan-epithelial VHL and HIF inactivation in the kidney to investigate renal epithelial HIF in the regulation of erythropoiesis. We found that acute HIF stabilization in renal epithelial cells resulted in rapid development of anemia and suppressed the induction of EPO under hypoxic conditions. This was associated with a shift of renal metabolism toward glycolysis, a reduction in mitochondrial mass and O₂ consumption, and an increase in cortical PtO₂ despite the presence of anemia. Our findings highlight what we believe is a novel role for renal epithelial HIF in erythropoiesis and suggest that oxygen-sensitive tubulointerstitial intercellular crosstalk modulates EPO production in REPCs.

Results

Pan-epithelial Vhl inactivation in the kidney results in rapid development of renal anemia. In order to study the effects of constitutive HIF stabilization in renal epithelial cells, we generated a mouse model of conditional Vhl inactivation in which a luciferase and Cre-recombinase transgene (LC-1) is under control of a tetracycline-sensitive bidirectional promoter, which is activated by a reverse tetracycline-responsive transactivator (rtTA) under control of paired box 8 (Pax8) gene regulatory elements (14). As previously reported, Pax8-rtTA-based Cre/loxP-mediated recombination targets renal epithelial cells along the entire nephron (Figure 1A) as well as a subset of hepatocytes (14). To achieve efficient and reproducible Vhl inactivation, 4-week-old Vhlft/fl Pax8-rtTA LC-1 mice and their Cre-negative littermates were treated with doxycycline for 2 weeks (hereafter referred to as P8; Vhl^{fl/fl} mutants and control mice). Phenotypic analysis was carried out within 2 weeks after completion of doxycycline treatment. Genomic PCR was used to confirm efficient recombination in the kidney and liver; recombination in other organs was not detected (Figure 1B and data not shown). Pax8-rtTA-mediated inactivation of Vhl resulted in HIF-1 α and HIF-2 α stabilization in the kidney (Figure 1B), and immunohistochemical analysis revealed strong nuclear HIF-1α staining in both cortical and medullary tubular epithelial cells, whereas HIF-2α was predominantly found in cortical nephron segments (Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI74997DS1). The nuclei of renal interstitial cells and glomeruli remained negative for both HIF-1 α and HIF-2 α , which is in line with the published expression pattern of *Pax8*-rtTA (14). In P8;Vhl^{fl/fl} livers, HIF-2α was detected in a subpopulation of hepatocytes (approximately 5% of all hepatocytes) (Supplemental Figure 1A). HIF-2α stabilization in hepatocytes was associated with polycythemia (hematocrit [Hct]: $65.7\% \pm 0.7\%$ in P8; Vhl^{fl/fl} mutants vs. $47.7\% \pm 1.2\%$ in littermate controls; n = 3 each) (Figure 1C), which is consistent with findings from our and other laboratories (15, 16). Hepatic Epo transcript levels were strongly increased in P8;Vhlfl/fl mutant mice, while renal Epo mRNA levels were suppressed, indicating that the kidneys of P8;Vhl^{N/I} mutant mice responded physiologically to the presence of polycythemia and that REPCs were not targeted by Pax8-rtTA-mediated Cre recombination (Figure 1C). In contrast to the Pax8-rtTA model, Vhl inactivation in renal epithelium as well as in nonendothelial interstitial cells using a Cre transgene driven by the Pax3 promoter (4) increased renal Epo mRNA levels substantially (Supplemental Figure 1B). Taken together, our genetic data support the concept that renal epithelial cells do not have the capacity to synthesize EPO under conditions of constitutive HIF activation (Figure 1D).

In order to understand the polycythemia-independent effects of renal epithelial Vhl inactivation on the kidney, we established a Pax8-rtTA-based conditional Vhl knockout model, in which the supraphysiologic EPO production in the liver was ablated. For this, we generated *Vhl/Epo* double-knockout mice (P8;*Vhl*^{fl/fl} $Epo^{fl/fl}$), which were homozygous for both the *Vhl*- and the *Epo*-2-lox alleles. In this model, both alleles recombined with equal efficiency in kidney and liver (Figure 2A). Because REPCs were not targeted and because EPO synthesis in Vhl-deficient hepatocytes was ablated, we hypothesized that erythropoiesis in P8;Vhl^{n/fl} Epofl/fl mutant mice would be normal. Surprisingly, P8;Vhlfl/fl Epofl/fl mutant mice developed significant anemia rapidly, within 2 weeks after doxycycline treatment was initiated (Hct: 28.8% ± 0.8% vs. $43.4\% \pm 0.6\%$; Hb: 8.7 ± 0.4 g/dl vs. 13.0 ± 0.1 g/dl; rbc: $6.2 \pm 0.2 \text{ M/µl}$ vs. $9.2 \pm 0.1 \text{ M/µl}$; n = 5 each) (Figure 2B). Reticulocyte counts and reticulocyte production index (RPI) were reduced, which indicated that anemia was hypoproliferative (Figure 2C). In contrast, erythropoiesis in P8;*Epo^{fl/fl}* single-knockout mice was not impaired, which is in line with our observation that REPCs were not targeted in the Pax8-rtTA-based genetic model and which established that anemia was the result of Vhl inactivation and not Epo inactivation in epithelial cells (Figure 2B). Renal Epo transcript levels were significantly reduced in P8;Vhl^{fl/fl} Epo^{fl/fl} mutant mice, indicating that P8;Vhl^{n/fl} Epo^{fl/fl} mutants developed anemia due to insufficient renal Epo transcription, whereas nonanemic P8;Epofl/fl mutants produced normal renal Epo transcript levels at baseline (Figure 2D) and had normal sEPO responses to phlebotomy (Supplemental Figure 4B). Taken together, our findings demonstrate that acute ablation of Vhl in renal epithelial cells suppresses EPO production in the kidney and results in the rapid development of renal anemia.

Since anemia in patients with CKD develops when renal function declines by at least 50% and glomerular filtration rate (GFR) is less than 60 ml/min (3), we examined whether rapid anemia development in P8;Vhl^{fl/fl} Epo^{fl/fl} mice was accompanied by decreased renal function. We measured blood urea nitrogen (BUN) levels and determined GFR by measuring FITC-inulin clearance. We furthermore evaluated renal blood flow (RBF) by 99mTc-MAG3 scintigraphy 2 weeks after completion of doxycycline treatment. We did not observe significant differences in either BUN levels $(43.25 \pm 1.8 \text{ mg/dl in P8}; Vhl^{fl/fl} Epo^{fl/fl} \text{ mutants vs. } 36.75$ \pm 4.6 mg/dl in controls, n = 4 each), GFR values (234.5 \pm 38.4 μ l/ min in P8; $Vhl^{fl/fl}$ Epofl/fl mutants vs. 218.0 ± 13.0 μ l/min in controls, n = 3 each), or RBF (0.36 ± 0.01 relative activity/s in P8; $Vhl^{fl/fl}Epo^{fl/fl}$ mice vs. 0.32 ± 0.04 relative activity/s in controls, n = 4 each) between P8;Vhl^{f/fl} Epo^{fl/fl} and control mice (Supplemental Figure 2A). Plasma Na⁺ and K⁺ levels were normal, indicating that sodi-

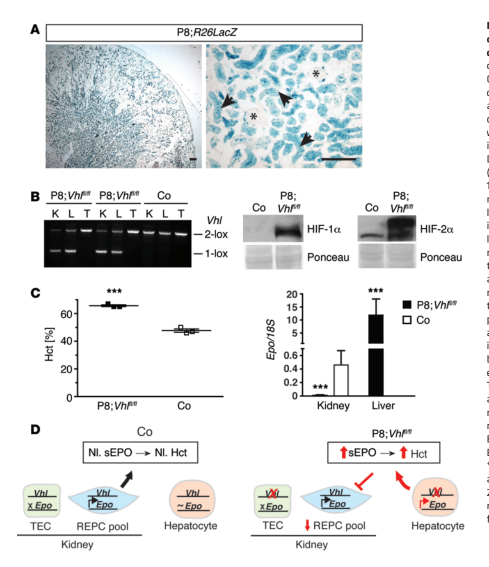


Figure 1. Pax8-rtTA-mediated inactivation of Vhl induces hepatic EPO and causes polycythemia. (A) Representative X-gal staining of the kidney from an R26R; Pax8-rtTA; LC-1 Cre-reporter mouse. Blue staining indicates cells that have undergone Pax8-rtTA-mediated Cre recombination. Cre activity was only detected in renal epithelial cells (arrows), whereas glomeruli were not stained (asterisks). (B) Left panel: PCR analysis of genomic DNA isolated from kidney (K), liver (L), and tail (T) of P8; VhI^{f1/f1} mutant and control (Co) mice 1 week after completion of doxycycline treatment. Right panel: HIF-1 α and HIF-2 α protein levels in whole-kidney lysates. Ponceau staining was used to demonstrate equal protein loading. (C) Hct values in individual P8;Vhlf^{fl/fl} mutant and control mice (n = 3 each) and relative Epo mRNA expression levels in kidneys and livers from P8;Vhlf1/f1 mutant and control mice (n = 3 each). (**D**) Schematic depicting the cellular contributions to sEPO under physiological conditions in control (left panel) and P8; $VhI^{fl/fl}$ mice (right panel). The red X indicates Cre-mediated gene inactivation. The black x indicates that the Epo gene in tubular epithelial cells (TEC) is not transcribed. The tilde indicates that Epo is nondetectable in hepatocytes at baseline and does not contribute to plasma EPO levels under normoxic conditions. In P8;Vhlf^{fl/fl} mutants, EPO is produced in hepatocytes, while renal EPO output is physiologically suppressed. ***P < 0.001, 2-tailed Student's t test. Shown are mean values \pm SEM. Scale bars: 200 μm . 2-lox, nonrecombined conditional allele; 1-lox, recombined allele. See Supplemental Figure 1 for additional information.

um and water homeostasis was grossly intact (Supplemental Figure 2B). Furthermore, morphologic and molecular examination of P8; $Vhl^{n/n}$ $Epo^{n/n}$ kidneys did not reveal any structural abnormalities, evidence of inflammation, or increased expression of renal injury markers, such as kidney injury molecule 1 (Kim1) and lipocalin-2 (Lcn2), compared with control (Supplemental Figure 2C). Thus, our data indicate that the rapid development of renal anemia in P8; $Vhl^{n/n}$ $Epo^{n/n}$ mice did not result from a loss of renal function or structural changes in the kidney and was not associated with renal inflammation.

Inactivation of epithelial Vhl reduces the number of REPCs. Previous studies have shown that the number of REPCs, which is dependent on the degree of renal hypoxia, controls renal EPO production (for a detailed overview see ref. 3). To examine whether acute Vhl ablation in renal epithelium affected the number of REPCs, we used high-resolution in situ hybridization to visualize REPCs in nonphlebotomized and phlebotomized P8; $Vhl^{0/ll}$ $Epo^{0/ll}$ and Crecontrol mice. We found that the number of REPCs at baseline was significantly reduced in P8; $Vhl^{0/ll}$ $Epo^{0/ll}$ mice compared with Crecontrol animals (1.7 \pm 0.8 cells/mm² in mutants and 9.8 \pm 0.5 cells/mm² in control mice). To examine EPO production in P8; $Vhl^{0/ll}$ $Epo^{0/ll}$ kidneys in response to anemia, we induced similar degrees

of anemia in P8; $Vhl^{N/l}$ $Epo^{N/l}$ and control mice by phelobotomy (Hct: 14.4% \pm 0.6% and 16.7% \pm 1.4%, respectively). We found that the predicted increase of REPC number was severely blunted in P8; $Vhl^{N/l}$ $Epo^{N/l}$ mice (37.1 \pm 12.4 cells/mm²) compared with control mice (210.2 \pm 19.1 cells/mm²) (Figure 3A). The decrease in REPC number was associated with a significant reduction in sEPO levels as well as a reduction in renal Epo transcript levels (Supplemental Figure 3A). In contrast with P8; $Vhl^{N/l}$ $Epo^{N/l}$ mutants, the level of renal Epo induction was similar between phlebotomized P8; $Epo^{N/l}$ and control mice, demonstrating that Pax8-rtTA-mediated inactivation of Epo alone in renal epithelium did not affect the physiologic behavior of renal interstitial cells with EPO-producing potential (Supplemental Figure 3B).

Renal cortical PtO₂ is elevated in P8;Vhl^{β / β} Epo^{β / β} mice. Because the size of the REPC pool is PtO₂ dependent, we next asked whether the reduction in REPC number in P8;Vhl^{β}/ β Epo^{β / β} mutants was associated with changes in renal PtO₂. Cortical PtO₂ was determined with a Clark electrode in 3 groups of mice: P8;Vhl^{β}/ β Epo^{β / β} mutants (Hct: 33.0% \pm 0.8%, obtained in 6 of 10 experimental mice), normemic control mice (Hct: 44.8% \pm 1.1%, obtained in 6 of 10 experimental mice), and phlebotomized anemic control mice (Hct: 20.1% \pm 1.5%, n = 7) (Supplemental Figure 3C). Despite

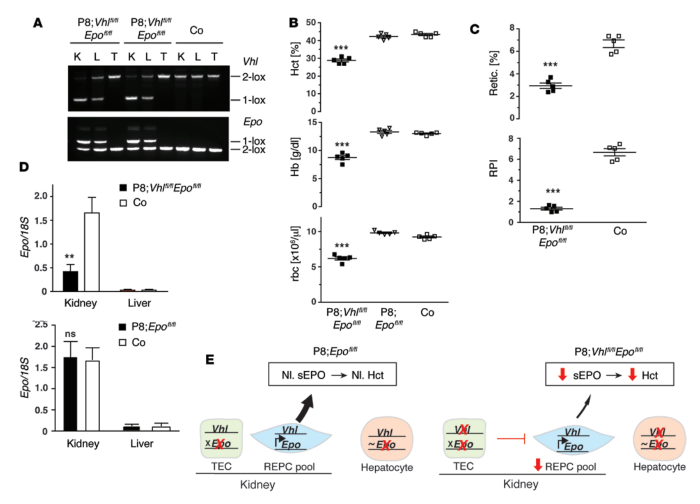


Figure 2. Renal epithelial *VhI* inactivation results in anemia. Hematologic effects of Pax8-rtTA-mediated inactivation of VhI in the absence of hepatic EPO induction. (**A**) Genomic PCR analysis of DNA isolated from kidney, liver, and tail of P8; $VhI^{IP/IP}$ $Epo^{IP/IP}$ and Control mice. (**B**) Hct, Hb, and rbc counts in P8; $VhI^{IP/IP}$ $Epo^{IP/IP}$, P8; $Epo^{IP/IP}$, and control (n = 5 each). (**C**) Corresponding reticulocyte counts (Retic.) and RPI in P8; $VhI^{IP/IP}$ $Epo^{IP/IP}$ and control mice (n = 5 each). (**D**) Relative renal and hepatic Epo mRNA levels in P8; $VhI^{IP/IP}$ $Epo^{IP/IP}$ and control mice (top panel, n = 5 each) and P8; $Epo^{IP/IP}$ (bottom panel; n = 6 kidney samples and n = 3 liver samples) in comparison with control mice (n = 8 kidney samples and n = 3 liver samples). (**E**) Schematic depicting cellular contributions to sEPO levels in P8; $Epo^{IP/IP}$ single mutants (left panel) and P8; $VhI^{IP/IP}$ $Epo^{IP/IP}$ double mutants (right panel). Red x indicates Cre-mediated gene inactivation; black x indicates that the Epo gene is not transcribed in tubular epithelial cells. The tildes indicate that Epo is not detectable in hepatocytes under baseline conditions. EPO production in the liver is no longer increased in P8; $VhI^{IP/IP}$ $Epo^{IP/IP}$ double mutants (right panel). **P < 0.01; ***P < 0.001, 1-way ANOVA followed by Dunnett's post hoc analysis (**B**); unpaired 2-tailed Student's t test (**C** and **D**). Shown are mean values t SEM. 2-lox, nonrecombined conditional t or t or t and t or t or t or t and t or t

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the presence of anemia, cortical PtO_2 in $P8;Vhl^{pl/l}$ $Epo^{n/l}$ mice was significantly elevated compared with that in normemic controls (39.26 \pm 2.2 mmHg vs. 30.8 \pm 1.3 mmHg; n = 10 for both groups; P < 0.01). PtO_2 values in anemic control mice averaged 14.2 \pm 2.6 mmHg, which was expected, as oxygen-carrying capacity in these mice was significantly reduced (n = 7) (Figure 3B). Arterial O_2 saturation (SaO_2) or PaO_2 was found to be indistinguishable between mutants and control mice, indicating that differences in arterial oxygenation did not account for the observed differences in cortical PtO_2 (Supplemental Figure 3C).

To gain additional insights into the regulation of EPO in P8; $Vhl^{ll/ll}$ $Epo^{ll/ll}$ mutants, we asked whether EPO suppression in P8; $Vhl^{ll/ll}$ $Epo^{ll/ll}$ mutants was exclusively dependent on PtO₂. Inhibition of PHD catalytic activity stabilizes HIF-2 α and increases EPO synthesis irrespective of PtO₂. Assuming equal bioavailability, we predicted that systemic administration of a potent pan-HIF

prolyl-4-hydroxylase inhibitor (PHI) would result in equally efficient HIF activation in kidneys from mutant and control mice and lead to comparable Epo transcription if the PHD/HIF/EPO axis was functionally intact in interstitial fibroblast-like cells. For the pharmacologic inhibition of HIF prolyl-4-hydroxylation, we utilized tool compound GSK1002083A (GlaxoSmithKline). We have previously shown that GSK1002083A resulted in robust activation of the HIF/EPO axis regardless of PtO, levels (4). While PHI treatment led to an approximately 16-fold increase in renal Epo mRNA levels in control mice, Epo induction in PHI-treated P8; Vhlf^{l/fl} Epo^{fl/fl} mutants was blunted (~7-fold increase compared with control; n = 8each; P < 0.001) (Figure 3C). This finding suggested that the PHD/ HIF/EPO axis in P8;Vhlft/ft Epoft/ft mutants was less sensitive to PHD inhibition and raised the possibility that anemia development in these mutants was not exclusively due to increased cortical PtO, and may also have involved PtO2-independent mechanisms.

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Epo^{fl/fl}

Phlebotomy

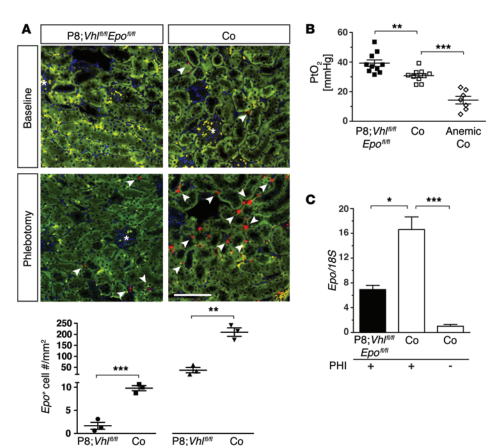


Figure 3. The number of EPO-producing cells is reduced in P8;Vhlfi/fi Epofi/fi mutants. (A) Representative images of Epo mRNA detection by RNA in situ hybridization (red spots) in kidney section from P8;Vhlfl/fl Epofl/fl and Cre-control mice at baseline and following phlebotomy. DAPI (blue) was used for nuclear staining. Number of Epo-expressing cells was counted in each renal cross-section and expressed as number of cells/mm² (n = 3 each). (B) Shown are the results from cortical tissue O, partial pressure (PtO₂) measurements in P8;Vhlf^{fl/fl} Epofi/fi and control mice. Each data point represents the average mean value of at least 3 measurements per individual mouse (n = 10, 10, and 7, respectively). (C) Relative Epo mRNA levels in wholekidney homogenates from P8;VhIf1/f1 Epofi/fi and control mice treated with PHI GSK1002083A 1 week after completion of doxycycline treatment (n = 8, 8, and3, respectively). *P < 0.05; **P < 0.01; ***P < 0.001, unpaired 2-tailed Student's t test (A); 1-way ANOVA followed by Dunnett's post-hoc analysis (B and C). Shown are mean values ± SEM. Scale bar: 100 μm. See Supplemental Figure 3 for additional information.

Vhl ablation in renal epithelium shifts renal metabolism toward glycolysis. We next asked whether changes in renal metabolism could be responsible for the increased cortical PtO2, as HIF regulates cellular energy metabolism by increasing glucose utilization and glycolysis and shifts oxidative phosphorylation toward glycolysis (17) by blocking the conversion of pyruvate to acetyl-CoA via inhibition of pyruvate dehydrogenase (PDH) (18, 19). To assess the effects of pan-tubular HIF activation on renal energy metabolism in vivo, we first performed microarraybased genome-wide mRNA expression analysis in whole-kidney homogenates from Vhl-deficient mice. We found that a substantial number of genes involved in glucose and fatty acid metabolism were differentially expressed (>1.5-fold change with an adjusted P value of < 0.05). The mRNA levels of most genes regulating glycolysis and glucose metabolism were increased (Figure 4). These included hexokinase 2 (Hk2), glucosephosphate isomerase 1 (Gpi1), aldolase C (Aldoc), triosephosphate isomerase 1 (Tpi1), phosphogylcerate kinase 1 (Pgk1), phosphoglycerate mutase 1 (Pgam1), pyruvate kinase M2 (Pkm2), and lactate dehydrogenase A (Ldha) (Figure 4A). Gene-expression changes were in support of our hypothesis that renal epithelial energy metabolism in Vhl-deficient kidneys had shifted from oxidative phosphorylation toward glycolysis. Consistent with a reduction in oxidative phosphorylation was the differential expression of genes involved in fatty acid storage and lipogenesis, which suggested a dysregulation of fatty acid metabolism (Figure 4B). In addition,

Epo^{fl/f}

we observed decreased expression of mRNAs encoding multiple ATP-consuming renal transporters, such as the sodium-dependent dicarboxylate transporter (Nadc1/Slc13a2), the sodium/sulfate symporter (NaS2/Slc13a4), sodium/hydrogen exchanger 2 (Nhe2/Slc9a2), the ammonium transporter rhesus blood groupassociated B glycoprotein (Rhbg/Slc42a2), the sodium-dependent neutral amino acid transporter (B[0]at1/Slc6a19), sodiumdependent phosphate transport protein 2B (Npt2b/Slc34a2), and the organic anion transporter 2 (Oat2/Slc22a7), raising the possibility that epithelial Vhl deletion led to a reduction in renal energy demand (Figure 4C). Microarray data were validated by realtime PCR analysis (Figure 4D). The metabolic gene-expression changes were furthermore reflected in increased glucose uptake in the renal medulla, as assessed by ¹⁴C-deoxy-D-glucose (2-DG) uptake measurements, and in elevated total kidney renal pyruvate and lactate levels, as measured by mass spectrometry in P8;Vhl^{f/fl} Epo^{f/fl} mice (Figure 4, E and F). Taken together, our data suggest that Vhl inactivation in renal epithelium led to marked changes in kidney metabolism and shifted renal energy metabolism from oxidative metabolism toward glycolysis, which in turn predicted a decrease in renal O₂ consumption.

Vhl ablation in renal epithelium decreases O_2 consumption. To assess renal O_2 consumption in Vhl-deficient kidneys, we performed high-resolution respirometry in mechanically permeabilized cortical and medullary tissue samples from P8;Vhl^{0/fl} and P8;Vhl^{0/fl} Epo^{fl/fl} mutant kidneys. We assessed respiratory activity

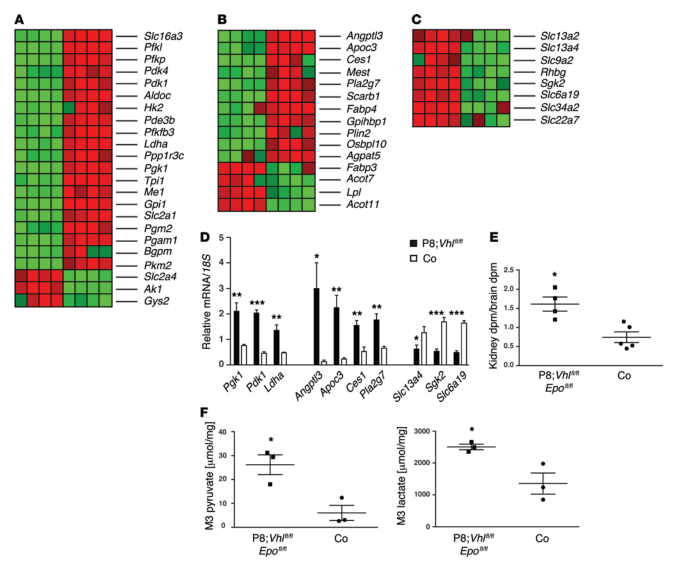


Figure 4. Epithelial Vhl ablation alters renal metabolism. Differential regulation of genes involved in glycolysis (**A**), fatty acid metabolism (**B**), and renal transport function (**C**) by genome-scale RNA expression profiling in kidneys from P8;Vhl^{fi/fj} and Cre[−] control mice (n = 4 each). Shown are differentially regulated metabolic genes that were either significantly up- or downregulated (≥ 1.5-fold). Color intensity is scaled within each row so that the highest expression value corresponds to bright red and the lowest to bright green. Each column represents expression levels in an individual kidney. (**D**) Verification of gene-expression analysis by quantitative PCR for key target genes from P8;Vhl^{fi/fj} and Cre[−] control mice (n = 6 each). (**E**) Assessment of renal glucose metabolism by quantification of 2-DG uptake in renal medulla from P8;Vhl^{fi/fj} Epo^{fi/fj} and control mice (n = 3 each). (**F**) Renal M3 pyruvate and M3 lactate concentrations in total kidney extracts from P8;Vhl^{fi/fj} Epo^{fi/fj} and control mice (n = 3 each). *P < 0.00; **P < 0.00; **P < 0.001, unpaired 2-tailed Student's t test. Shown are mean values t SEM.

at complex I (cI), complex II (cII), and complex IV (cIV), utilizing complex-specific substrates and inhibitors. In cortical and medullary tissue preparations, state 2 respiration (lack of ADP in the presence of substrate) and the activities of cI, cII, and cIV were markedly reduced in both P8; $Vhl^{p/n}$ and P8; $Vhl^{p/n}$ $Epo^{n/p}$ kidneys (shown here are data for P8; $Vhl^{p/n}$ $Epo^{n/p}$ and control kidneys; n=5, respectively) (Figure 5A). Because HIF regulates mitochondrial biogenesis (2O), we next assessed mitochondrial mass in Vhl-deficient mice. For this, we measured the activity of mitochondrial citrate synthase (CS) in cortical and medullary tissue samples (21). Both cortical and medullary CS activity, which typically correlate with mitochondrial mass, were significantly reduced, by approximately 30% and 50%, respectively. This finding was furthermore reflected in a comparable reduction of mitochondrial DNA (mtDNA)

(Figure 5, B and C). Single mitochondrial respiration in cortex, which was assessed by normalizing O_2 consumption to CS activity, was not different between P8; $Vhl^{p/p}$ $Epo^{p/p}$ and control mice (n=5 each). Normalized O_2 consumption in medullary preparations was also unchanged for state 2, cI, and cII respiration, while single mitochondrial O_2 consumption through cIV was increased (Figure 5D). In summary, our results demonstrate that reduced renal O_2 consumption in Vhl-deficient mice was associated with a reduction in mitochondrial mass.

Cocontribution of epithelial HIF-1 α and HIF-2 α to anemia development in P8;Vhl^{N/I} Epo^{N/I} mice. Because both HIF-1 α and HIF-2 α were detected in renal tissue from Vhl-deficient mice (Figure 1), we generated P8;Vhl^{N/I} Epo^{N/I} Hif1a^{N/I} and P8;Vhl^{N/I} Epo^{N/I} Hif2a^{N/I} triple-knockout mice to characterize the individual contributions of HIF-1 α

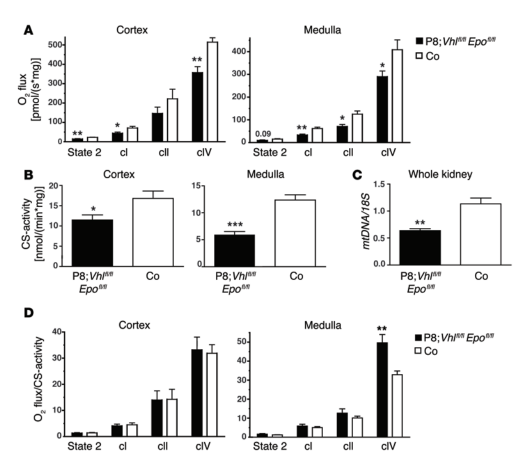


Figure 5. Renal epithelial Vhl ablation reduces respiratory capacity and mitochondrial mass. (A) O, flux measurements in mechanically permeabilized cortical and medullary tissue samples from P8; VhIf1/f1 Epof1/f1 and Cre^{-} control mice (n = 5each). (B) CS activity of cortical and medullary tissue samples from P8; $VhI^{fl/fl}$ $Epo^{fl/fl}$ and control mice (n = 5 each). (\mathbf{C}) Relative quantification of mitochondrial mass by real-time PCR analysis of whole-kidney genomic DNA samples from P8;VhIf1/f1 Epof1/f1 and control mice (n = 5 each). Relative mtDNA levels were normalized to nuclear encoded 185 DNA. (D) O, fluxes normalized to CS activity (n = 5 each). *P <0.05; **P < 0.01; ***P < 0.001, unpaired 2-tailed Student's t test. Shown are mean values ± SEM. State 2, basal respiration in the presence of substrate but lack of ADP.

and HIF-2 α to the development of anemia and metabolic reprogramming in P8; $Vhl^{N/\beta}$ $Epo^{N/\beta}$ double mutants. Genomic PCR analysis was used to confirm that efficient recombination of all conditional alleles $(Vhl^{N/\beta}, Epo^{N/\beta}, \text{ and } Hifla^{N/\beta} \text{ or } Hifla^{N/\beta})$ was achieved, and only mice with efficient recombination of the respective alleles in renal tissue were included in the analysis (Supplemental Figure 4A). Complete blood count analysis showed that erythropoiesis remained impaired in both P8; $Vhl^{N/\beta}$ $Epo^{N/\beta}$ $Hifla^{N/\beta}$ and P8; $Vhl^{N/\beta}$ $Epo^{N/\beta}$ $Hifla^{N/\beta}$ mutants (Hct: 30.0% \pm 1.3%; Hb: 9.2 \pm 0.3 g/dl; rbc: 6.4 \pm 0.2 M/µl; n = 6 and Hct: 34.3% \pm 0.9%; Hb: 10.7 \pm 0.2 g/dl; rbc: 7.9 \pm 0.2 M/µl; n = 3) (Figure 6A), suggesting that both HIF- α homologs contributed to the development of anemia in P8; $Vhl^{N/\beta}$ $Epo^{N/\beta}$ mice.

Although renal *Epo* transcript levels in P8;*Vhl*^{II/I} *Epo*^{II/I} *Hif1a*^{II/II} and P8;*Vhl*^{II/I} *Epo*^{II/II} *Hif2a*^{II/II} mice were statistically not significantly different from those of control animals (Figure 6A), they were inadequate for the degree of anemia found in triple-mutant mice. Anemia of nonrenal etiology is usually characterized by an inverse linear relationship between Hct and the decadic logarithm of sEPO concentration. This relationship is altered in anemia associated with CKD, as the diseased kidney fails to respond to hypoxic stimuli with adequate EPO production (22). sEPO levels in P8;*Vhl*^{II/II} *Epo*^{II/II}, P8;*Vhl*^{II/II} *Epo*^{II/II}, and P8;*Vhl*^{II/II} *Epo*^{II/II} mutant mice were inappropriately low for the degree of anemia that was present, whereas a linear decrease in Hct was associated with the expected logarithmic increase in sEPO levels in both P8;*Epo*^{II/II} and control mice (Supplemental Figure 4B).

To investigate whether HIF-1 and HIF-2 served redundant functions with regard to the observed changes in metabolic gene

expression in P8; $Vhl^{IV,II}$ $Epo^{IV,II}$ mice, we measured expression levels of genes involved in glycolysis (Pgk1 and pyruvate dehydrogenase kinase 1 [Pdk1]), lipid metabolism (angiopoietin like 3 [Angpt13] and apolipoprotein C-III [Apoc3]), and transport functions (Slc13a4 and serum/glucocorticoid regulated kinase 2 [Sgk2]). As shown in Figure 6B, upregulation of glycolytic gene expression appeared to be exclusively HIF-1 dependent, whereas the expression of Angpt13 and Apoc3 appeared to be HIF-2 dependent. While we could not attribute the suppression of transport genes Slc13a4 and Sgk2 to a specific HIF- α homolog, coinactivation of either Hif1a or Hif2a in P8; $Vhl^{IV,II}$ $Epo^{II,II}$ mice was sufficient to increase mitochondrial mass (Figure 6C). Taken together, our genetic data suggest that the development of anemia in P8; $Vhl^{IV,II}$ $Epo^{IV,II}$ mice required both HIF-1 and HIF-2 activation and could not be attributed to one HIF homolog alone.

Regulation of EPO synthesis by tubular epithelial cells is HIF dependent and localizes to the proximal nephron. Because we dissected the role of HIF-1 and HIF-2 in the development of anemia in Vhl-deficient mice and found that inactivation of either homolog did not restore normal erythropoiesis, we investigated to what degree anemia development in P8;Vhl^{P_i/P_i} Epo^{P_i/P_i} mice required intact signaling through both HIF- α homologs. For this, we bred mice that carried a floxed Hif1b (Arnt) allele to P8;Vhl^{P_i/P_i} Epo^{P_i/P_i} mice, generating P8;Vhl^{P_i/P_i} Epo^{P_i/P_i} Arnt^{P_i/P_i} triple-knockout mice. ARNT serves as the β -subunit for both HIF-1 and HIF-2 heterodimers, and its inactivation abolishes canonical HIF signaling in renal epithelium (23). Concomitant ablation of Arnt restored hematologic parameters to normal values, as no significant differ-

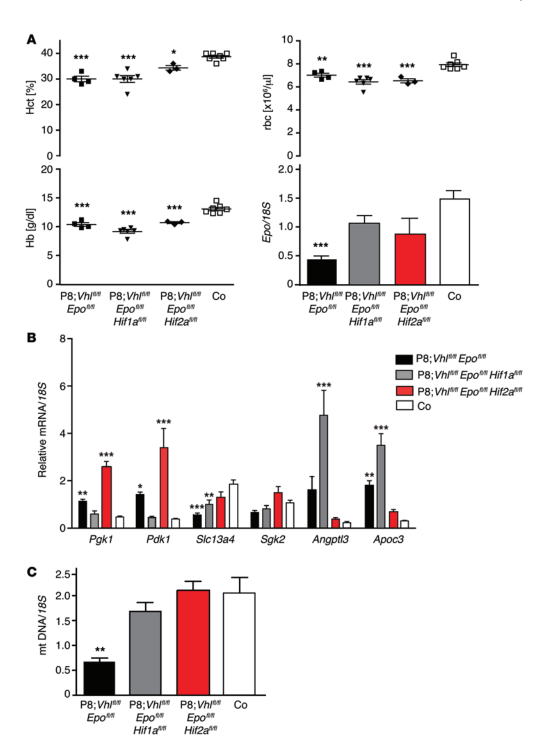


Figure 6. Epithelial HIF-1 and HIF-2 coregulate renal Epo production and metabolism. (A) Shown are Hct, Hb, and rbc counts as well as renal Epo mRNA levels in P8;Vhlfl/fl $Epo^{fl/fl}$, P8; $Vhl^{fl/fl}$ $Hif1a^{fl/fl}$, P8; $Vhl^{fl/fl}$ Hif2af1/f1, and Cre-control mice (n = 4, 6, 3, and 7, respectively).(B) Renal mRNA expression of metabolic genes involved in glucose metabolism, fatty acid metabolism, and renal transport in P8; VhIf1/f1 Epof1/f1, P8; VhIf1/f1 Hif1afl/fl, P8;VhIfl/fl Hif2afl/fl, and Cre $^-$ control mice (n = 4, 6, 3,and 7, respectively). (C) Relative quantification of mitochondrial mass by real-time PCR analysis of wholekidney genomic DNA samples from P8; VhIf1/f1 Epof1/f1, P8; VhIf1/f1 Hif1afl/fl, P8;Vhlfl/fl Hif2afl/fl, and Cre $^-$ control mice (n = 4, 6, 3,and 7, respectively). *P < 0.05; **P < 0.01; ***P < 0.001, 1-way ANOVA followed by Dunnett's post hoc analysis (A and C); unpaired 2-tailed Student's t test (B). Shown are arithmetic mean values ± SEM. See Supplemental Figure 4 for additional information.

ences in Hct, Hb, or rbc were found between P8; $Vhl^{N/l}$ Epo^{N/l} Arnt^{N/l} and control mice (Figure 7A). This finding is consistent with renal *Epo* transcript levels in P8; $Vhl^{N/l}$ Epo^{N/l} Arnt^{N/l} mice that were comparable to those of control mice (Figure 7A). These genetic data establish that the suppression of renal EPO production in P8; $Vhl^{N/l}$ Epo^{N/l} mice required intact HIF signaling.

To investigate which specific segment of the nephron is involved in the suppression of renal EPO production, we examined several conditional *Vhl* knockout lines in which different nephron segments were targeted (*Pepck-Cre*, renal proximal tubule; *Thp-Cre*, medullary thick ascending limb of Henle and early distal

tubule; Hoxb7-Cre, collecting duct). Since Pepck-Cre-driven conditional ablation of Vhl leads to polycythemia due to induction of hepatic EPO production (23), we generated $Pepck-Cre\ Vhl^{N/R}\ Epo^{R/R}$ double-knockout mice. Similar to the P8; $Vhl^{R/R}\ Epo^{R/R}$ mutants, $Pepck-Cre\ Vhl^{R/R}\ Epo^{R/R}$ double-knockout mice were anemic (6 to 8 weeks old, Hct: $29.4\%\pm1.5\%$; Hb: 6.8 ± 0.4 g/dl; rbc: $7.1\pm0.3\times10^6/\mu$ l; n=6), whereas mice with heterozygous deletion of Vhl and homozygous deletion of $Epo\ (Pepck-Cre\ Vhl^{R/R}\ Epo^{R/R})$ displayed normal erythropoiesis (Hct: $39.0\%\pm0.7\%$; Hb: 12.9 ± 0.2 g/dl; rbc: $8.8\pm0.1\times10^6/\mu$ l; n=4), (Figure 7B). Hematological analysis of 6- to 8-week-old $Thp-Cre\ Vhl^{R/R}\ (n=5)$ and $Hoxb7-Cre\ Vhl^{R/R}$

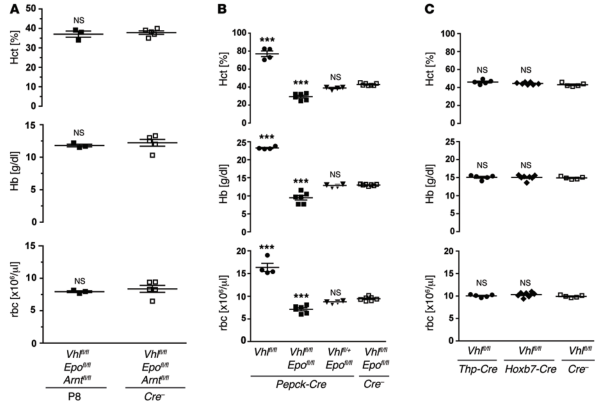


Figure 7. The renal proximal tubule regulates EPO production in an HIF-dependent manner. (A) Upper panels show normal Hct, Hb, and rbc counts in P8; $Vh^{fl/fl}$ $Epo^{fl/fl}$ $Arnt^{fl/fl}$ mutant mice and Crecontrol mice (n=4-5). Lower panel shows relative Epo mRNA expression levels in kidney homogenates from P8; $Vh^{fl/fl}$ $Epo^{fl/fl}$, P8; $Vh^{fl/fl}$ $Epo^{fl/fl}$, and control mice (n=4,4, and 7, respectively). (B) Shown are Hct, Hb, and rbc counts for individual 6- to 8-week-old $Pepck-Cre\ Vhf^{fl/fl}$, $Pepck-Cre\ Vhf^{fl/fl}$, $Epof^{fl/fl}$, and Crecontrol mice $Epof^{fl/fl}$, $Epof^{fl/fl}$, $Epof^{fl/fl}$, and $Epof^{fl/fl}$, $Epof^{fl/fl}$,

(n=7) mice revealed no differences in Hct, Hb, or rbc values compared with Cre-negative littermate controls (n=5) (Figure 7C), suggesting that Vhl ablation in proximal renal epithelium alone is sufficient for the suppression of renal EPO production. In summary, our data establish that the development of anemia in P8; $Vhl^{p/p}$ mice was HIF dependent and suggest that HIF activation via Vhl gene deletion in the proximal nephron modulates renal EPO production (Figure 8).

P8;Vhf

Epoten Arntini

Со

Discussion

2.0

Epo/18S

0.5

0

P8;Vhl™

Epo^{ft/fl}

In the current study, we utilized inducible and cell type-specific Cre recombinase-mediated gene targeting to investigate the role of non-EPO-synthesizing cell types in the regulation of EPO production in the kidney. We demonstrate that HIF activation in renal epithelial cells via acute Vhl gene deletion suppressed erythropoiesis by restricting the number of interstitial fibroblast-like cells that are able to convert to REPCs. This effect localized to the proximal nephron and was associated with renal gene expression and functional changes that indicated a shift from oxidative phosphorylation toward glycolysis as well as decreased O_2 consumption due to reduced mitochondrial mass. As a result of epithelial HIF acti-

vation, renal cortical ${\rm PtO_2}$ was elevated despite the presence of marked anemia. Our data suggest that HIF-regulated tubuloint-erstitial intercellular crosstalk suppresses EPO production under conditions of sustained HIF activation in the kidney. We propose a model whereby HIF activation in a renal cell type that does not synthesize EPO modulates renal EPO production by regulating the number of REPCs.

Ascent to high altitude triggers a pronounced systemic hypoxia response, which results in increased renal EPO synthesis and rbc production. Interestingly, sEPO levels decline rapidly before significant changes in Hct are noted (12, 13). While the reasons for this phenomenon are not clear, increased sequestration of EPO from serum by EPO-expressing cells in the bone marrow has been proposed as a possible underlying mechanism (2). The decline in sEPO levels is, however, paralleled by a reduction in renal *Epo* mRNA expression pointing toward intercellular crosstalk that is directed at local EPO synthesis (24). While it is possible that the hypoxic induction of PHD3 might reduce HIF levels and suppress *Epo* transcription in REPCs, there is currently no experimental evidence that PHD3 alone regulates renal EPO synthesis (25–29). Findings from our genetic studies, however, would lend experimental sup-

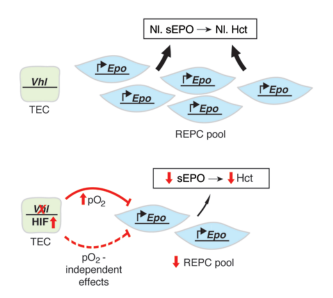


Figure 8. Renal epithelial cells regulate erythropoiesis. Model depicting the role of renal tubular epithelial cells in the regulation of EPO production at baseline (upper panel) and under conditions of constitutive HIF activation as a result of *VhI* inactivation (lower panel). Constitutive HIF stabilization in renal tubular epithelial cells inhibits the conversion of non-EPO-producing peritubular interstitial cells to REPCs.

port to the notion that HIF-regulated metabolic adaptation of renal tubular epithelial cells to hypoxia suppresses EPO synthesis in the kidney partly via modulation of renal O₂ consumption.

In our model, both epithelial HIF-1 and HIF-2 were involved in modulating renal EPO production at baseline and under conditions of anemic hypoxia, as concomitant deletion of either Hifla or Hif2a in P8;Vhlf1/f1 Epof1/f1 mice did not restore erythropoiesis to control levels and resulted in inadequate sEPO levels for the degree of anemia observed in these mutants. While epithelial HIF activation in P8;Vhl^{fl/fl} Epo^{fl/fl} mice was associated with anemia, we did not find that inactivation of either epithelial HIF-1α or HIF-2α alone (P8;Hif1a^{fl/fl}, P8;Hif2a^{fl/fl}) or together (P8;Hif1a^{fl/fl} Hif2a^{fl/fl}) enhanced renal EPO responses when mice were exposed to prolonged hypoxia (8% O, for 6 days) or phlebotomy (data not shown). This may be due to the degree of hypoxia generated in this model, i.e., levels of HIF- α stabilization, hypoxic adaptation over time, or timing of the analysis. Our model of nonhypoxic HIF activation is based on Vhl gene deletion in renal epithelial cells alone, which leads to the complete loss of the ability to degrade HIF- α and would mimic severe hypoxia. Under severe hypoxia, factor inhibiting HIF (FIH), an HIF asparagine hydroxylase that hydroxylates an asparagine residue in the C-terminal transactivation domain of HIF- α , would further modify HIF-mediated hypoxia responses, as it operates as a second hypoxic switch that regulates recruitment of coactivators to the HIF transcriptional complex (5). Nevertheless, our model provides genetic evidence for HIF-regulated tubulointerstitial cell interaction affecting the conversion of interstitial cells into EPO-producing cells. It is plausible that in the hypoxic kidney, additional nonepithelial cell-derived paracrine signals that also modulate the renal EPO response may come into play.

EPO synthesis occurs in peritubular interstitial fibroblast-like cells, which are normally found in the inner cortex and corticome-

dullary region (7, 30). The main stimulus for renal EPO production is hypoxia, and small changes in renal oxygenation are rapidly translated into increased EPO synthesis. PtO, is therefore a critical determinant of renal EPO output. The regulation of renal PtO, is complex and depends largely on arterial O2 content, on GFR-driven tubular work load, which in turn determines O, consumption, and on the degree of arteriovenous O, shunting, which is a specific feature of the renal microenvironment that results from the counter current exchange of O₂ between arterial to venous vessels that are arranged in parallel (31, 32). Since GFR and RBF change in parallel under most conditions, the boost in arterial O, delivery that normally associates with a rise in RBF is largely offset by an increase in O2 consumption, thus limiting its impact on PtO2. As a result, regional PtO, stays within a relatively narrow range and is highly susceptible to changes in intrarenal metabolism (3, 31). HIF activation is known to have profound effects on cellular metabolism. This occurs at multiple levels and includes the increased expression of glucose transporters and glycolytic enzymes, the regulation of TCA cycle flux, and fatty acid oxidation and storage as well as mitochondrial biogenesis, respiration, and autophagy, which could explain the rapid decrease in mitochondrial mass and O, consumption in our model (17).

We used genetic means to further dissect the contribution of individual nephron segments to the regulation of erythropoiesis and identified the proximal tubule as the nephron segment that modulates EPO production in the kidney. Our observations are in line with previously published physiologic studies that used acetazolamide to block tubular sodium reabsorption in the rat renal proximal tubule. This resulted in a dose-dependent reduction of sEPO in response to hypoxia, possibly, although not measured in this report, through a decrease in renal $\rm O_2$ consumption (11). Consistent with these findings are also studies in humans, which demonstrated that sEPO levels diminish when climbers, who ascend to high altitude, are treated with acetazolamide (33).

The effects of epithelial HIF activation on erythropoiesis were already clearly detectable by day 10 of doxycycline treatment (data not shown). When considering the average mouse erythrocyte life span of approximately 40 days (34), the reduction in rbc numbers by approximately 35% within 2 weeks after initiation of doxycycline treatment indicates that the suppression of renal Epo transcription must have occurred rapidly. While the timing of EPO suppression is consistent with functional changes in renal metabolism that alter PtO2, our findings also indicate that PtO2independent mechanisms may have participated in the modulation of renal EPO synthesis, as the renal *Epo* response in P8;Vhl^{fl/fl} Epo^{fl/fl} mice remained blunted in response to systemic pharmacologic HIF prolyl-4-hydroxylase inhibition, which was predicted to robustly inhibit HIF PHDs and induce Epo in peritubular interstitial fibroblast-like cells irrespective of PtO2. The mechanisms underlying this observation are unclear and warrant further investigation.

Since our model of pan-epithelial HIF activation is based on *Vhl* gene deletion, it is likely that tubulointerstitial cellular crosstalk comes into play during the pathogenesis of VHL-associated renal cell cancer (RCC) and negatively affects the ability of surrounding peritubular interstitial fibroblast-like cells to synthesize EPO. In contrast with our model of pan-epithelial *Vhl* inactivation, these effects, however, may not become clinically apparent in

human RCC patients, as tumorigenesis occurs in a localized fashion and involves only a small fraction of renal epithelial cells.

Renal EPO output correlates with the number of REPCs (6, 7), i.e., the degree of decrease in PtO, determines the number of peritubular interstitial fibroblast-like cells that switch from a non-EPO-producing to an EPO-producing state. This kind of functional plasticity is not uncommon in the kidney and has been observed in other renal cell types. For example, afferent arteriolar perivascular cells change their functional phenotype and begin to produce renin in response to sodium depletion (35). While PtO, is a major determinant of REPC pool size, the regulation of REPC plasticity, especially the quality of intra- and extracellular signals that control the ability of interstitial fibroblast-like cells to switch to an EPOproducing state, is only incompletely understood. Myofibroblast transdifferentiation of renal interstitial fibroblast-like cells, which is associated with the development of renal fibrosis and increased extracellular matrix production, has been suggested as suppressing their EPO-producing potential, possibly through activation of NF-κB signaling (36). Here, we demonstrate that HIF activation in non-EPO-producing renal epithelial cells modulates REPC pool size and therefore may affect erythropoiesis under kidney injury conditions. This may be particularly important in chronic injury conditions where epithelial HIF stabilization has been shown to associate with the development of fibrosis (37).

Aside from renal interstitial cells, multiple other cell types have been shown to be capable of synthesizing EPO or of modulating erythropoiesis under certain experimental conditions (3). HIF stabilization induced by conditional ablation of *Vhl* or *Phds* in hepatocytes, astrocytes, renin-producing cells, osteoblasts, and other cell types led to increased sEPO levels and polycythemia through direct activation of tissue-specific *Epo* transcription (3). In contrast, HIF stabilization in non-EPO-producing keratinocytes causes polycythemia indirectly by redirecting blood flow to the skin, away from kidney and liver, which induced hypoxia and stimulated renal and hepatic EPO production (38). Our data indicate that renal epithelial HIF modulates EPO synthesis in peritubular interstitial fibroblast-like cells, thus providing evidence for kidney-specific non-cell autonomous control of EPO synthesis and suppression of EPO production in the context of HIF activation.

In summary, our genetic studies demonstrate that HIF activation in proximal tubular epithelial cells, which do not synthesize EPO, negatively modulates erythropoiesis. While more experimental studies are needed to further explore the functional relationships between epithelial cells and interstitial fibroblast-like cells in the kidney, our data provide a molecular basis for the participation of renal epithelial cells in the regulation of interstitial cell plasticity and pathogenesis of renal anemia.

Methods

Generation of mice and genotyping. The generation and genotyping of Vhl, Epo, Arnt, Hifla, and Hifla conditional alleles and the lacZ ROSA26 reporter (R26R) mouse line as well as LC-1, Pax8-rtTA, Pepck-Cre, Thp-Cre, Hoxb7-Cre, and Pax3-Cre transgenic lines have been described elsewhere (14, 39–47).

Inducible Cre-mediated renal tubular ablation of *Vhl*, *Arnt*, *Hifla*, *Hif2a*, and/or *Epo* was achieved by generating mice that carried both the rtTA-dependent Cre-recombinase (LC-1) transgene and

expressed rtTA under control of the *Pax8* promoter (*Pax8*-rtTA). The following genotypes were generated: (a) *Vhl^{N/R} Pax8*-rtTA LC-1, (b) *Epo^{R/R} Pax8*-rtTA LC-1, (c) *Vhl^{N/R} Epo^{R/R} Pax8*-rtTA LC-1, (d) *Vhl^{N/R} Epo^{R/R} Epo^{R/R} Hifla^{R/R} Pax8*-rtTA LC-1, and (f) *Vhl^{N/R} Epo^{R/R} Hif2a^{R/R} Pax8*-rtTA LC-1. After completion of doxycycline treatment, these were referred to as P8;*Vhl^{N/R}*, P8;*Epo^{R/R}*, P8;*Vhl^{N/R} Epo^{R/R} Arnt^{R/R}*, P8;*Vhl^{N/R} Epo^{R/R}*, P8;*Vhl^{N/R}*, P8;*Vhl^{R/R}*, and P8;*Vhl^{R/R}*, Epo^{R/R} *Hif2a^{R/R}*, respectively. For the activation of the *Pax8*-rtTA/LC1 conditional knockout system, 4-week-old mice were treated for 2 weeks with doxycycline (Sigma-Aldrich) contained in drinking water at a concentration of 0.2 mg/ml in 5% sucrose water.

Constitutive conditional inactivation of *Vhl* and/or *Epo* in specific nephron segments was achieved by generating mice that were homozygous for the *Vhl* and/or *Epo* conditional alleles and expressed either a proximal tubule–specific Cre transgene (*Pepck-Cre*), a distal tubule–specific Cre transgene (*Thp-Cre*), or a collecting duct–specific Cre transgene (*Hoxb7-Cre*). Analysis of nephron segment-specific knockout mice was performed at age 6 to 8 weeks. Renal interstitial fibroblast-like cells and tubular epithelial cells were targeted with the *Pax3-Cre* transgene. The following genotypes were generated: (a) *Pepck-Cre Vhl^{Il/II}*, (b) *Pepck-Cre Vhl^{Il/II}*, (c) *Pepck-Cre Vhl^{Il/II}*, (d) *Thp-Cre Vhl^{Il/II}*, (e) *Hoxb7-Cre Vhl^{Il/II}*, and (f) *Pax3-Cre Vhl^{Il/II}*, respectively.

DNA, RNA, and protein analysis. DNA and RNA isolation were performed as described previously (48). For real-time PCR analysis, 1 µl of cDNA was subjected to PCR amplification on an ABI StepOnePlus platform using either SYBR green or TaqMan Universal PCR Master Mix (Applied Biosystems). Relative mRNA expression levels were quantified with the relative standard curve method according to the manufacturer's instructions (Applied Biosystems). Primer sequences for the analysis of Epo, Kim-1, Lcn2, F4/80, and Tnfa expression have been published elsewhere (48, 49). For the analysis of Il1b expression, the following primers were used: fwd, 5'-GGTCAAAGGTTTG-GAAGCAG-3'; rev, 5'-TGTGAAATGCCACCTTTTGA-3'. All genes were normalized to 18S levels using the 18S TaqMan set from Applied Biosystems. Microarray analysis was carried out using an Affymetrix Exon/Gene (WT) microarray chip. Intensity values from each hybridization were scaled versus a global average signal from the same array and normalized by robust multichip average (RMA) analysis. Statistical analysis was performed using a moderated t test and Bonferroni's familywise error rate (FWER) multiple testing. Differentially regulated genes were defined by a 1.5-fold difference in mean expression values and an adjusted P value of less than 0.05 between groups. Heat maps were created using Heatmap Builder (http://ashleylab.stanford.edu/tools/tools-scripts.html). All original microarray data were deposited in the NCBI's Gene Expression Omnibus (GEO GSE54172). For RNA in situ detection using the RNAscope Muliplex Fluorescent Assay Kit, paraffin sections of mouse kidneys (5-µm thickness) were processed according to the manufacturer's protocol (Advanced Cell Diagnostics). The hybridized sections were imaged with the Ariol SL-50 automated slide scanner. Nuclear protein extracts for Western blot analysis were prepared as previously described; HIF-1α was detected with a polyclonal antibody from Novus Biologicals (catalog NB100-449), and HIF-2α was detected with a custom-made polyclonal antibody from Invitrogen, as previously described (15).

Pharmacological HIF activation. For pharmacological HIF activation, prolyl-4-hydroxylase inhibitor GSK1002083A (GlaxoSmith-

Kline) was dissolved in 1% methylcellulose and administered by oral gavage at a dose of 60 mg/kg twice, 24 and 4 hours prior to tissue harvest (4). Methylcellulose without compound served as control.

Blood work and assessment of renal function. Hcts were determined by microcapillary tube centrifugation in a Micro Hematocrit Centrifuge (International Equipment Co.) or with a Hemavet 950 analyzer (Drew Scientific). sEPO levels were determined by ELISA (R&D Systems). Reticulocyte counts were measured by FACS analysis of whole blood stained with thiazole orange (Sigma-Aldrich). BUN levels were determined using the QuantiChrom Urea Assay Kit (BioAssay Systems), and serum Na⁺ and K⁺ concentrations were determined with an iStat analyzer and EC8⁺ cartridges (Abaxis). For determination of SaO₂ and PaO₂, blood samples were analyzed with iStat analyzer and CG8⁺ cartridges (Abaxis). GFR was assessed using the FITC-inulin single-injection method as previously described (50). GFR was calculated based on a 2-compartment model using GraphPad Prism (GraphPad Software). RBF was estimated by ^{99m}Tc-MAG3 dynamic scintigraphy as previously described (51).

Histological analysis. For HIF-1α and HIF-2α immunohistochemistry, polyclonal rabbit anti-HIF-1α antibody (Cayman Chemical, catalog 10006421) and polyclonal rabbit anti-HIF-2α anti-serum PM8 (52) were used at dilutions of 1:10,000 and 1:20,000, respectively, and detected with the CSA-II High Signal Amplification Kit and the Rabbit Link Reagent (Dako). For detection of CD45 $^{+}$ leukocytes, anti-mouse monoclonal anti-CD45 antibody (BD Biosciences — Pharmingen, catalog 550539) was used and detected with the Vectastain Elite Kit (Vector). X-gal staining was performed on frozen tissues sections according to MacGregor (53).

High-resolution respirometry. Respiratory activities in tissue samples from cortex and medulla of mutant and control mice were analyzed with the O2k Oxygraph System (Oroboros). Tissue samples from the cortex and medulla were dissected, minced on ice, and placed on ice in mitochondrial respiration buffer (final concentration: 110 mM sucrose, 60 mM K-lactobionate, 0.5 mM EGTA, 1 g/l BSA essentially fatty acid free, 3 mM MgCl2, 20 mM taurine, 10 mM KH2PO4, 20 mM Hepes, adjusted to pH 7.1 with KOH) at a concentration of 1 mg/ml. Samples from P8;Vhlf^{t/fl} Epo^{fl/fl} and control mice were placed into an O2k Oxygraph (Oroboros), and O, consumption and flux were determined for state 2 (presence of substrate in the absence of ADP), cI, cII, and cIV separately by titration of the following substances: (a) glutamate/ malate (final concentration: 10/2 mM), (b) ADP (final concentration: 25 mM), (c) rotenone (final concentration: 0.5 μM), (d) succinate (final concentration: 10 mM), (e) antimycin A (final concentration: 2.5 μM), and (f) TMPD/ascorbate (final concentration: 0.5/2 mM). Measurements were live recorded using DatLab software (Oroboros).

Metabolic analysis and cortical PtO₂ measurements. CS activity was measured as previously described (21). Briefly, aliquots from samples used for high-resolution respirometry were homogenized, and 20 μl of homogenate was added to 180 μl medium containing 0.1 mM 5,5-dithio-bis-(2-nitrobenzoic) acid (DTNB) (Sigma-Aldrich), 0.5 mM oxaloacetate (Sigma-Aldrich), 50 M EDTA (Sigma-Aldrich), 0.31 mM acetyl coenzyme A (Sigma-Aldrich), 5 mM triethanolamine hydrochloride (Fluka), and 0.1 M Tris-HCl, pH 8.1 (Sigma-Aldrich). The activity of CS was measured spectrophotometrically at 412 nm. Glucose uptake was measured in renal medulla with 2-DG at 13 μCi/mouse and normalized to brain as previously described (54). Polar metabolites were analyzed as previously described (55). Mice were fasted for 4 hours prior to intravenous

injection of U-13C glucose (20 mg/mouse). At 25 minutes following the injection, mice were sacrificed and kidneys were harvested. ¹³C isotopic enrichment was analyzed in total kidneys using Agilent 6890/5975N GC-MS equipped with 25 m DB-5ms capillary column for analysis of isotopic enrichment.

For cortical ${\rm PtO}_2$ measurements, mice were subjected to inhalation anesthesia with isoflurane. Core body temperature was monitored with a rectal probe and maintained between 36.5°C and 37.5°C. The left kidney was exposed through flank incision and was stabilized without causing excessive strain on renal vessels. The kidney capsule was removed, and the left kidney was covered with mineral oil. Renal ${\rm PtO}_2$ was measured with a modified Clark-type microelectrode (\emptyset = 8–10 $\mu {\rm m}$, Unisense). Two-point calibration was carried out in water saturated with either ${\rm N}_2$ gas or ambient air at 37°C. For every mouse, a minimum of 3 independent measurements were obtained per kidney 1 mm below the renal surface after a linear correlation between ${\rm O}_2$ tension and electric current was established. For statistical analysis, only average mean values from individual mice were used.

Statistics. Data reported represent mean values \pm SEM. Statistical analyses were performed with Prism 5.0b software (GraphPad Software). When 2 groups of mice were compared, the unpaired, 2-tailed Student's t test was used; for comparison of 3 and more groups, 1-way ANOVA followed by Dunnett's post hoc analysis was used. P < 0.05 was considered statistically significant.

Study approval. All procedures involving mice were performed in accordance with NIH guidelines for the use and care of live animals and were approved by the Institutional Animal Care and Use Committee (IACUC) of Vanderbilt University.

Author contributions

NMF and VHH conceived and designed the research studies, analyzed and interpreted data, wrote the manuscript, and created the figures. NMF, QL, HK, FS, and OD performed experiments and acquired and analyzed data. TAI, PMO, and JF helped with design and interpretation of research studies or contributed reagents or technical expertise.

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