U19/Eaf2 Binds to and Stabilizes von Hippel-Lindau Protein

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Abstract

Studies have firmly established a key regulatory role for the tumor suppressor pVHL in the regulation of the vascular system and normal spermatogenesis. Here, we report that knockout of the newly identified tumor suppressor U19/Eaf2 also caused vascular system abnormalities and aspermatogenesis, suggesting a potential link between U19/Eaf2 and pVHL. Immunoprecipitation and in vitro binding assays showed an association between U19/Eaf2 and pVHL, whereas deletion mutagenesis revealed the requirement of the NH2 terminus of U19/Eaf2 and both the α and β domains of pVHL for this binding. U19/Eaf2 stabilizes pVHL, as shown by protein stability and pulse-chase studies. Testes and mouse embryonic fibroblasts (MEF) derived from U19/Eaf2 knockout mice expressed reduced levels of pVHL, indicating that full in vivo expression of pVHL indeed requires U19/Eaf2. As expected, U19/Eaf2 knockout MEF cells exhibited an increased level and activity of hypoxia-inducible factor 1α (HIF1α), a protein typically regulated via a pVHL-mediated degradation pathway. Furthermore, angiogenesis in a Matrigel plug assay was significantly increased in U19/Eaf2 knockout mice. The above observations argue that U19/Eaf2 can modulate HIF1α and angiogenesis, possibly via direct binding and stabilization of pVHL. [Cancer Res 2009;69(6):2599–606]

Introduction

Carriers of germ-line mutations in the von Hippel-Lindau (VHL) gene are predisposed to the VHL syndrome. This autosomal dominant syndrome has multiple manifestations, including high incidence of clear cell renal carcinoma, hemangioblastomas, and pheochromocytomas (1–4). Conventional VHL deletion in the mouse is embryonic lethal due to vascular abnormalities in the placenta (5). However, mice with conditional VHL deletions survive, and studies using these mice further support a tumor-suppressive role for pVHL (6, 7). Targeted deletion of VHL in the livers of BALB/c mice resulted in the hypervascularization of the liver parenchyma (6). Mice with tissue-specific VHL deletions also developed hepatic vascular tumors, angiectasis in multiple organs, and defects in spermatogenesis (7), pointing to the essential role of pVHL in the homeostasis of multiple tissues.

pVHL contributes to the regulation of oxygen-responsive gene expression by targeting the transcription factor hypoxia-inducible factor 1α (HIF1α) for proteasomal degradation under normoxic conditions (8, 9). In a mechanism analogous to the SCF(Skp1/Cdc53/F-box) ubiquitination machinery, pVHL targets HIF1α for degradation by serving as a bridge between HIF1α and a multiprotein E3 ubiquitin ligase complex consisting of elongin C, elongin B, Cul-2, and Rbx 1 (10–13). In this capacity, pVHL acts in a manner similar to an F-box protein (9). pVHL has two main domains: α domain (residues 155–192) and β domain (residues 163–154 and 193–204; ref. 14). The α domain binds to the E3 ubiquitin ligase complex via elongin C (15, 16), whereas the β domain contains the substrate recognition motif for HIF1α. Association of pVHL with HIF1α requires the hydroxylation of key proline residues (P402 and P564) within the oxygen degradation domain of HIF1α. Under normoxic or atmospheric O2 levels (21%), a conserved family of prolyl-4-hydroxylases mediates hydroxylation of HIF1α, allowing pVHL to bind to and thus inhibit HIF1α for ubiquitin-mediated proteasomal degradation (17–22). In contrast, limited O2 inhibits proline hydroxylation, preventing the association between pVHL and HIF1α. Consequently, HIF1α accumulates and enters the nucleus, where it dimerizes with the constitutively stable HIF1β subunit to form the heterodimeric transcription factor HIF (20). As an active transcription factor, HIF acts as the primary regulator of both cellular and systemic responses to decreased O2 concentration; responses include enhanced glycolysis, cell proliferation, apoptosis, erythropoiesis, and angiogenesis (23).

Loss of pVHL, therefore, leads to the constitutive activation of HIF1α and its target genes, mimicking a state of constant hypoxia despite normal O2 content. The inappropriately expressed HIF1α targets include vascular endothelial growth factor (VEGF), platelet-derived growth factor B, and mitogenic factors such as transforming growth factor α (24–26). Overproduction of the proangiogenic factors may explain the hypervascular nature of hemangioblastomas and renal cell carcinomas. Although HIF is a critical target of pVHL in the progression of hemangioblastomas and renal cell carcinoma, pVHL may also interact with other proteins critical to tumor suppression (27–29). Thus, the identification and characterization of additional pVHL binding partners will help elucidate the mechanism of pVHL-mediated tumor suppression and/or the regulation of pVHL function.

U19/Eaf2 is a newly identified tumor suppressor. We previously reported evidence for U19/Eaf2 down-regulation, allelic loss, promoter hypermethylation, and homozygous deletion in human advanced prostate cancer specimens. U19/Eaf2 is also implicated in acute myeloid leukemia. Consistent with its potential suppressive role in the human, U19/Eaf2 knockout mice developed lung adenocarcinoma, hepatocellular carcinoma, B-cell lymphoma, and high-grade prostatic intraepithelial neoplasia (30).

Here, we report that the U19/Eaf2 knockout mice displayed abnormalities in vasculogenesis and spermatogenesis. VHL conditional knockout mice also developed abnormalities in vasculogenesis and spermatogenesis, suggesting a potential relationship between U19/Eaf2 and pVHL. Indeed, we present evidence that U19/Eaf2 can bind to...
and stabilize pVHL, thus blocking HIF-driven angiogenesis. Our data argue that U19/Eaf2 can modulate the pVHL pathway via direct binding and stabilization of pVHL.

**Materials and Methods**

**Histologic analysis.** Analyses were done using virgin male mice of ages 1 to 24 mo. Experimental cohorts were wild-type, heterozygous, and homozygous littermates with C57BL/6 and 129 combined genetic backgrounds. Tissues were fixed in 10% phosphate-buffered formalin at 4°C. Samples were then embedded in paraffin, sectioned at 5 μm, and stained with H&E. Immunohistochemical staining was done as described previously (30).

**Cell culture and transfection.** RCC4, 293, and Cos-7 cells were cultured in DMEM with 10% fetal bovine serum (FBS) and penicillin/streptomycin. RCC4/vec and RCC4/VHL (provided by Peter J. Ratcliffe, University of Oxford, Oxford, United Kingdom) were cultured in DMEM with 10% FBS, 1 mg/mL G418, 0.1 mmol/L nonessential amino acids (Invitrogen), 100 μmol/L 2-mercaptoethanol (Sigma), and penicillin/streptomycin. Primary mouse embryonic fibroblasts (MEF) were immortalized by SV40T antigen (provided by Katherine Rundell, Northwestern University, Chicago, IL) and maintained in DMEM with 10% FBS, 0.1 mmol/L nonessential amino acids (Invitrogen), 100 μmol/L 2-mercaptoethanol (Sigma), and penicillin/streptomycin. Cos-7 and MEFs were transfected using Lipofectamine 2000 (Invitrogen). 293 cells were transfected using Fugene 6 (Roche).

**Antibodies and chemical reagents.** Anti-Flag (M2; monoclonal) and anti-c-Myc–conjugated agarose were purchased from Sigma. Monoclonal anti-hemagglutinin (HA) antibody was purchased from Covance Research Products. Anti-c-Myc (polyclonal), antivmouse VHL (polyclonal), and anti-HA–conjugated agarose were purchased from Santa Cruz Biotechnology; anti-c-Myc (polyclonal), and antihuman VHL (polyclonal) were purchased from BD Biosciences. Anti-HIF1α (NB-100-449) for mouse HIF1α detection was purchased from Novus Biologicals. Cycloheximide was purchased from Sigma.

**Plasmids.** We subcloned Myc-tagged pVHL, U19/Eaf2, and their mutants into pcMV-Myc vector (Clontech) at SalI and KpnI sites by PCR. Wild-type pcDNA VHL was a gift from Peter J. Ratcliffe. Flag-tagged VHL was a gift from Frank Lee (Belfast City Hospital, Belfast, United Kingdom). The hypoxia response element (HRE) and mutated HRE-luciferase reporter constructs were provided by Navdeep Chandel (Northwestern University, Chicago, IL). To obtain the construct for pVHL in vitro translation, we subcloned pVHL into pBluescript II KS+ vector (Stratagene). Full-length pVHL and U19/Eaf2 cDNA were subcloned into pG3H3AM vector (31) to generate HA-tagged proteins (a methionine-rich variant provided by William Tansey, Cold Spring Harbor Laboratory, New York, NY).

**Immunoprecipitation and Western blotting.** Wild-type Flag-pVHL and Myc-U19/Eaf2 or Myc empty vector were transiently transfected into Cos-7 cells. Wild-type Myc-U19/Eaf2 and HA-pVHL or HA empty vector were transiently transfected into 293 cells. After 24 h, cells were directly lysed in radioimmunoprecipitation assay (RIPA) buffer with complete protease inhibitors (Roche). For coimmunoprecipitation, we incubated 500 μg of whole-cell extract with 20 μL of Myc-agarose conjugate (Sigma) or 10 μL of HA-agarose conjugate (Santa Cruz Biotechnology) for 16 h at 4°C. The immune complex was washed three times in five times in RIPA buffer, separated by SDS-PAGE (NuPAGE 4–12% Bis-Tris Gel, Invitrogen), and analyzed by Western blot.

**pVHL-expressing stable revertants of RCC4** (RCC4/VHL) were maintained by hypoxia (1% O2) for 16 h to enhance U19/Eaf2 protein levels and lysed in RIPA buffer. For immunoprecipitation, we incubated 500 μg of whole-cell extract for 1 h with 20 μL of protein A/G-agarose beads (Santa Cruz Biotechnology) and 1 μg of normal mouse immunoglobulin, which were then incubated with 50 μL of a rabbit polyclonal antibody against human U19/Eaf2 or 1 μg of normal rabbit IgG for 16 h at 4°C. The immune complexes were washed in RIPA buffer, separated by SDS-PAGE, and analyzed by Western blot.

**In vitro association experiment.** Glutathione S-transferase (GST) and GST-U19/Eaf2 were purified from bacterial lysates using glutathione-agarose (Sigma). 35S-labeled pVHL was translated in vitro using a TNT-coupled reticulocyte lysate system (Promega). After preincubation in 0.1% bovine serum albumin/PBS for 1 h, GST or GST-U19/Eaf2 bound to glutathione-agarose and 35S-labeled pVHL were incubated at 4°C for 4 h. The beads were extensively washed five times with PBS containing 0.5% NP-40 and 150 mmol/L NaCl before being subjected to SDS-PAGE followed by autoradiography.

**Protein stabilization and pulse-chase experiment.** We transiently transfected Cos-7 and 293 cells with 2 μg of Flag-VHL or HA-VHL with increasing amounts of Myc-U19 or Myc-U19 (114–260) complemented by up to 2 μg total of Myc empty vector. After 28 h, the transfected cells were directly lysed in RIPA buffer, separated by SDS-PAGE, and analyzed by Western blotting.

We transiently transfected 293 cells with HA-VHL and either Myc-U19 or Myc-U19 (114–260). After 24 h, cells were treated with cycloheximide at 50 μg/mL for 0, 2, 4, and 8 h (32). Then, the cells were directly lysed in RIPA buffer, separated by SDS-PAGE, and analyzed by Western blot.

For pulse-chase assays, HA-VHL, and Myc-U19 or Myc empty vector were transiently transfected into 293 cells. After 48 h, cells were maintained in DMEM/10% FBS lacking Met and Cys for 1 h. Cells were then incubated in DMEM/10% FBS containing a [%E/Met/Cys mixture (50 μCi/mL; Perkin-Elmer) for 2 h. The cells were then washed twice with warm PBS and labeling was stopped by addition of growth medium with Met and Cys (3 mmol/L).

**Hypoxia treatment and luciferase assay.** U19+/+ and U19−− MEFs were maintained in a hypoxia chamber (1% O2) for 16 h to achieve HIF1α stabilization and then released into normoxia. Lysates were collected at multiple time points for immunodetection of HIF1α. Cycloheximide was used to prevent any new synthesis of HIF1α during normoxia.

We transiently transfected HRE and mutated HRE-luciferase reporter constructs along with the Renilla luciferase reporter into MEFs of U19+/+ and U19−−. After 6 h, the cells were placed in hypoxic conditions or maintained in normoxia for 20 h and then directly lysed using PLB buffer supplied by Dual Luciferase Reporter Assay system (Promega). The luciferase activity was determined using a luminometer (Siris, Zylux Corp.) and normalized to Renilla luciferase activity. Data are reported as mean ± SE of two separate experiments done in triplicate.

**Matrigel plug angiogenesis assay.** Age-matched U19/Eaf2 wild-type and U19/Eaf2-null mice (~6 mo old) were injected with 0.5 mL Matrigel (BD Biosciences) containing 2 ng/mL VEGF and 69 units/mL heparin. Three females and three male mice were used for each group. After 12 d, Matrigel plugs were removed for frozen sectioning. Matrigel sections (5 μm) were incubated with rat anti-mouse CD31 antibody (Pharmingen) followed by rhodamine-conjugated donkey anti-rat antibody (Jackson Immunoresearch). Digital images were obtained with a Nikon TE2000-U fluorescence microscope and quantified for microvascular density with ImageJ software.

**Results**

**U19/Eaf2 gene deletion causes late-onset aspermatogenes and extradermal hematoepoiiesis.** In our recently generated U19/Eaf2 knockout mice, we had observed cardiac enlargement, lung adenocarcinoma, hepatocellular carcinoma, B-cell lymphoma, and high-grade prostatic intraepithelial neoplasia (30). The mice also developed noncancerous phenotypes, such as aspermatogenesis seen in older animals (Fig. 1A–C). Human, mouse, and rat testes all express U19/Eaf2 (data not shown), suggesting a possible role in testicular function. Within 2 to 3 months of age, no reproduction abnormalities were detected in U19/Eaf2-null males or females (data not shown). However, at 22 months or older, 26.1% (12 of 46) of U19/Eaf2-null male mice exhibited a dramatic reduction in their testis size and wet weight; testes from normal animals weighed ~60% less (P < 0.05; Fig. 1A and C). H&E staining revealed that these abnormal testes contained degenerated germ...
cells filled with vacuoles; multinucleated giant cells, reflecting abnormal sperm maturation; and few Sertoli cells (Fig. 1A). Cross sections of the U19/Eaf2 knockout testes showed seminiferous tubule atrophy and collapse of the testicular capsule, with few sperm in the seminiferous tubule. Terminal deoxyribonucleotidyl transferase–mediated dUTP nick end labeling (TUNEL) staining revealed large numbers of germ cells undergoing apoptosis in U19/Eaf2-null males (Fig. 1A). These data indicate an important role of U19/Eaf2 in normal spermatogenesis during adulthood. Moreover, no testicular abnormality was observed in a cohort of 15 U19/Eaf2 heterozygotes and 20 wild-type mice (data not shown), suggesting that one copy of U19/Eaf2 is sufficient for normal development and maintenance of the testis.

We observed that two of the U19/Eaf2 knockout mice developed extramedullary hematopoiesis, accompanied by extreme enlargement of the spleen and liver (Fig. 1D). Histologic examination revealed spleen tissue filled with hematopoietic cells and immature blood cells at different stages, cells that normally exist in the bone marrow. More Ki-67–positive cells were present in the spleen of U19/Eaf2-null mice as compared with wild-type controls (Fig. 1D). Given the high expression level of U19/Eaf2 in bone marrow (33), U19/Eaf2 inactivation may disrupt the normal hematopoiesis in bone marrow, causing a compensatory response from spleen and liver. We recognize the possibility that an unrelated sporadic mutation may cause this phenotype because only 2 of 46 U19/Eaf2-null mice developed extramedullary hematopoiesis. However, we have never observed this phenotype in all of the wild-type or U19/Eaf2 heterozygotes in our lab, suggesting that the U19/Eaf2 knockout is likely responsible.

U19/Eaf2 protein binds to pVHL directly. Our studies showed that U19/Eaf2-null mice displayed defects in spermatogenesis (Fig. 1A) and in the vascular system, including heart enlargement (30) and extramedullary hematopoiesis (Fig. 1D). Work from others had established that conditional knockout of VHL, a well-characterized tumor suppressor in renal cell carcinoma, also causes defects in spermatogenesis and vascular system (7). Together, these observations led us to speculate that a functional link may exist between U19/Eaf2 and the VHL pathway. We first analyzed the possibility of a physical interaction between U19/Eaf2 and pVHL proteins by transiently coexpressing Myc-tagged U19/Eaf2 with Flag-tagged pVHL in either monkey kidney Cos-7 cells or human kidney 293 cells. Immunoprecipitation followed by Western blot analysis showed that Flag-tagged pVHL coprecipitated with Myc-tagged U19/Eaf2 with Flag-tagged pVHL in either monkey kidney Cos-7 cells or human kidney 293 cells. Immunoprecipitation followed by Western blot analysis showed that Flag-pVHL coprecipitated with Myc-U19/Eaf2, but not in vector-transfected controls (Fig. 2A). Similarly, Myc-U19/Eaf2 coprecipitated with HA-tagged pVHL (HA-pVHL; Fig. 2B), pointing to a specific association between exogenous U19/Eaf2 and pVHL. The reciprocal coimmunoprecipitation was done in both cell lines and was reproducible (data not shown), indicating that the cell lines used did not affect the association.
To test whether pVHL and endogenous U19/Eaf2 proteins interact, we performed coimmunoprecipitation experiments using RCC4 renal cancer cells expressing HA-tagged wild-type pVHL. We incubated cells under hypoxic conditions (1% O2) for 16 hours before the cell extract preparation because we have found that this increases U19/Eaf2 protein levels (data not shown). Antibody to U19/Eaf2 coprecipitated HA-pVHL, whereas isotype control IgG did not (Fig. 2C), arguing for an in vivo association between U19/Eaf2 and pVHL.

To assess whether U19/Eaf2 binds to pVHL directly, we performed in vitro binding assays using bacterially expressed and purified GST-U19/Eaf2 and in vitro translated 35S-labeled pVHL. Figure 2D

Figure 2. Interaction of U19/Eaf2 with pVHL. A, Flag-tagged pVHL was cotransfected with Myc-tagged U19/Eaf2 or empty Myc vector into Cos-7 cells. Cell lysates were immunoprecipitated (IP) with anti-Myc antibody–conjugated agarose beads, and blots were probed with anti-Flag antibody (top image) or anti-Myc antibody (bottom image). B, Myc-U19 was cotransfected with HA-pVHL or empty HA vector into 293 cells. Cell lysates were immunoprecipitated with anti-HA antibody–conjugated agarose beads, and blots were probed with anti-Myc antibody (top image) or anti-HA antibody (bottom image). C, whole-cell lysates from RCC4/VHL cells exposed to hypoxia (1% O2) for 16 h were subjected to immunoprecipitation with control rabbit IgG or rabbit anti-U19/Eaf2 polyclonal antibody, separated by SDS-PAGE, and immunoblotted with anti-HA and anti-U19/Eaf2 antibodies. D, in vitro translated 35S-labeled pVHL was incubated with bacterially purified GST-U19/Eaf2 fusion protein attached to glutathione-agarose beads. GST served as negative control. After incubation, the beads were extensively washed and subjected to SDS-PAGE. Association of 35S-VHL and U19/Eaf2 in vitro was detected by autoradiography. Bottom image, inputs of GST and GST fusion proteins are shown as Coomassie blue–stained gel.

Figure 3. Domain mapping of the interaction between U19/Eaf2 and pVHL. A, pVHL interacted with the NH2-terminal region of U19/Eaf2 in coimmunoprecipitation assays. HA-pVHL was cotransfected with Myc-tagged U19/Eaf2 COOH-terminal deletion mutants (1–67, 1–113, 1–161, and 1–245) and NH2-terminal deletion mutants (36–260, 68–260, 114–260, and 162–260) into 293 cells. Wild-type Myc-U19/Eaf2 was cotransfected with HA empty vector as a negative control. Cell lysates were immunoprecipitated with anti-HA antibody–conjugated agarose beads, and blots were probed with anti-Myc antibody (top image) or anti-HA antibody (bottom image). B, both the α and β domains of pVHL were required for interaction with U19/Eaf2 in coimmunoprecipitation assays. HA-U19/Eaf2 was cotransfected with the Myc-tagged major β domain (1–154), α and major β domains (1–192), α and entire β domains, α domain deletion mutant (Δ155–192), or major β domain deletion mutant (Δ64–154) into 293 cells. Wild-type Myc-pVHL cotransfected with HA empty vector served as negative control. Cell lysates were immunoprecipitated with anti-HA antibody–conjugated agarose beads, and blots were probed with anti-Myc antibody (top image) or anti-HA antibody (bottom image).
shows that pVHL bound GST-U19/Eaf2, but not GST alone, indicating a direct association between wild-type U19/Eaf2 and pVHL.

To further characterize the interaction between U19/Eaf2 and pVHL, we carried out domain mapping for both U19/Eaf2 and pVHL. We constructed eight different deletion mutants of U19/Eaf2 and tested their ability to bind pVHL in vivo (Fig. 3A). This analysis revealed that pVHL bound to the NH2 terminus of U19/Eaf2 (residues 1–113) but not to the COOH terminus (residues 114–260; Fig. 3A). We also constructed five deletion mutants of pVHL for use in coimmunoprecipitation experiments (Fig. 3B). pVHL required both its α and β domains to bind U19/Eaf2. Furthermore, the VHL mutant (residues 1–192) lacking the minor β domain (residues 193–204) displayed a decrease in U19/Eaf2 binding as compared with the VHL mutant (residues 1–206) containing the entire α and β domains. These data show that both the α and β domains are required for pVHL to bind U19/Eaf2.

U19/Eaf2 binding causes an increase in pVHL stability. While studying U19/Eaf2-pVHL interactions, we observed an increase in Flag-pVHL levels when it was cotransfected with Myc-U19/Eaf2 or Myc empty vector into 293 cells. Forty-eight hours after transfection, cells were metabolically labeled with [35S]Met/Cys for 2 h followed by pulse-chase analysis of Myc-U19/Eaf2 or Myc vector into 293 cells. Twenty hours after transfection, cells were treated with cycloheximide (CHX) at 50 μg/mL for the indicated number of hours. C, pulse-chase assay to determine HA-VHL protein half-life in the absence or presence of Myc-U19/Eaf2. Two micrograms of HA-VHL expression vector were cotransfected with 2.0 μg of Flag-VHL in the absence (Fig. 4A) or presence of increasing amounts of Myc-U19/Eaf2 (114–260), which does not bind to pVHL (Fig. 4A). These results indicate that U19 stabilizes pVHL and identify the NH2-terminal region (1–113) of U19 as critical for this stabilization.

We further evaluated the effect of U19/Eaf2 on pVHL stability using the protein synthesis inhibitor cycloheximide and pulse-chase experiments. We transfected HA-pVHL expression vector with Myc-U19/Eaf2, empty vector, or Myc-U19/Eaf2 (114–260) into 293 cells and, 24 hours later, treated cells with cycloheximide. Using Western blot analysis, we showed that in the presence of wild-type U19/Eaf2, the estimated half-life of HA-pVHL was longer than 8 hours (Fig. 4B). In contrast, when cells were cotransfected with empty vector control or the deletion mutant Myc-U19/Eaf2 (114–260), the estimated half-life of pVHL was no greater than 2 and 4 hours, respectively (Fig. 4B). The pulse-chase analysis further confirmed the ability of U19/Eaf2 to prolong the half-life of pVHL.
pVHL. (Fig. 4C). The above observations show that U19/Eaf2 enhances pVHL stability.

**U19/Eaf2 knockout tests have reduced levels of pVHL.** To determine if the endogenous pVHL level is decreased in the U19/Eaf2 knockout mice, we performed Western blot analysis with an anti-pVHL antibody. Given that the aspermatogenesis phenotype was observed in both VHL conditional knockout and U19/Eaf2-null mice, we initially measured VHL protein levels in the testes of U19/Eaf2 wild-type and U19/Eaf2-null mice. As expected, endogenous expression of pVHL was higher in the wild-type than in the U19/Eaf2-null testes (Fig. 4D). The endogenous pVHL protein levels in other major organs, including kidney, lung, and heart, were also reduced in the U19/Eaf2-null mice; however, the reduction was not as dramatic as that in testes (data not shown). Furthermore, endogenous pVHL levels in the U19/Eaf2-null MEF cells were lower than those in the wild-type MEF cells (Fig. 4D). To test whether U19/Eaf2 affects pVHL transcription, we carried out real-time reverse transcription-PCR analysis for VHL mRNA in 293 cells overexpressing U19/Eaf2 or control vector, testes from U19/Eaf2-null and wild-type mice, and U19/+/+ and U19/−/− MEF cells. These experiments revealed no difference in the VHL transcript levels in the matched sample pairs (data not shown). These data indicate that U19/Eaf2 influences pVHL level posttranslationally by stabilizing pVHL protein through direct binding.

**U19/Eaf2 deletion enhances HIF1α level and angiogenesis.** A well-documented function of pVHL is the targeting of HIF for oxygen-dependent proteolysis (8, 9). Given our finding that U19/Eaf2 modulates the degradation of HIF1α in immortalized U19+/+ and U19/Eaf2 had any effect on the pVHL-mediated degradation of HIF1α in immortalized U19+/+ and U19/−/− MEFs. As expected, the endogenous HIF1α level was higher in U19/+/+ cells than in U19/−/− cells, particularly under hypoxic conditions (Fig. 5A). When wild-type MEF cells were switched from hypoxia to normoxia, the majority of HIF1α was degraded within 5 minutes. In contrast, in U19/Eaf2-null MEF cells switched from hypoxia to normoxia, HIF1α was only partially degraded at 5 minutes or even 10 minutes after the switch. These results argue that U19/Eaf2 modulates the degradation of HIF1α, presumably mediated by pVHL.

We next examined the transcriptional activity of HIF1α in U19+/+ and U19/−/− MEF cells using a HRE-driven luciferase reporter (34). The HIF1α transcriptional activities in both U19/Eaf2-null and wild-type MEF cells were markedly induced under hypoxic conditions (Fig. 5B). As expected, the levels of HIF1α transcriptional activation in U19/−/− MEF cells were higher than those in U19+/+ cells, indicating a role of U19/Eaf2 in regulating HIF1α function. As a control, we showed that a mutant HRE construct with deleted HIF binding sites did not respond to the hypoxic induction of transcriptional activity in both U19+/+ and U19/−/− MEF cells.

Because U19/Eaf2 mediated efficient degradation of HIF1α, we next tested whether U19/Eaf2 could influence angiogenesis in the Matrigel plug assay (35, 36). Immunostaining with anti-CD31 antibody followed by quantitative analysis showed that the microvessel density in Matrigel plugs containing 2 ng/mL VEGF was ~3.5-fold higher in the U19/Eaf2-null mice relative to the wild-type mice, and the difference was statistically significant (P < 0.05; Fig. 6). This finding argues that U19/Eaf2 plays a role in regulating angiogenesis in cooperation with pVHL via the HIF1α pathway.

**Discussion**

We recently identified U19/Eaf2 as a tumor suppressor in multiple mouse tissues and possibly also in humans (30, 33). In this article, we show that U19/Eaf2-null mice develop noncancerous phenotypes that overlap with those observed in VHL conditional knockout mice. Furthermore, we present evidence for physical and functional interactions between U19/Eaf2 and pVHL, a well-established tumor suppressor.

The U19/Eaf2 knockout mice have diverse phenotypes in different tissues. We previously reported that U19/Eaf2 can function as a tumor suppressor as evidenced by the development of B-cell lymphoma, hepatocellular carcinoma, lung adenocarcinoma, and prostatic intraepithelial neoplasia in U19/Eaf2 knockout mice (30). Here, we show that U19/Eaf2 deletion also caused various nontumorigenic abnormalities, including aspermatogenesis and extramedullary hematopoiesis (Fig. 1). The diverse phenotypes observed in U19/Eaf2 knockout mice indicate that U19/Eaf2 could influence multiple cellular activities in various tissues and that U19/Eaf2 function seems to be cell type specific. In testes, U19/Eaf2 deletion significantly enhanced apoptosis but had no effect on proliferation. In contrast, U19/Eaf2 deletion enhanced proliferation but did not affect cell death in other organs such as the spleen. It is also interesting that the penetrance of U19/Eaf2-null mutation to develop each of the different phenotypes is not 100%. For example, the penetrance for the aspermatogenesis phenotype by U19/Eaf2 knockout is only ~26%.

![Figure 5](https://example.com/figure5.png)
U19/Eaf2 may regulate diverse cellular activities by binding to different partners. It was previously reported that U19/Eaf2 binds to ELL family proteins and thereby affects ELL transcription elongation activities. The report that Eaf1, the homologue of U19/Eaf2, is a component of the Cajal body, a nuclear suborganelle involved in the biogenesis of small nuclear ribonucleoproteins, suggests that U19/Eaf2 may also contribute to the formation of small nuclear ribonucleoproteins (37). Identification and characterization of additional U19/Eaf2 binding partners will provide further insights into the mechanisms by which U19/Eaf2 regulates various cellular processes.

Identification of pVHL as a new U19/Eaf2 binding partner implicates a role for U19/Eaf2 in pVHL signaling, providing a novel mechanism in U19/Eaf2 action. The interaction between U19/Eaf2 and pVHL seems to be physiologically relevant. The cycloheximide inhibition and pulse-chase experiments provide strong evidence that U19/Eaf2 can prolong the half-life of pVHL in cultured cells (Fig. 4). The observations that MEF cells, testes, and other tissues derived from U19/Eaf2-null cells had less pVHL relative to controls argue that U19/Eaf2 indeed stabilizes pVHL in vivo (Fig. 4). Furthermore, several pieces of evidence indicate that the interaction has functional consequences. For example, U19/Eaf2-null cells had higher HIF1α protein levels than wild-type controls, particularly under hypoxic conditions (Fig. 5). The prompt and complete degradation of HIF1α following release from hypoxia in wild-type MEF cells was retarded in the U19/Eaf2 knockout MEF cells. In addition, U19/Eaf2 deletion significantly enhanced HRE activation. These findings argue that U19/Eaf2 loss results in the elevation of the VHL/HIF/HRE axis, a pathway known to control angiogenesis (38). As expected, Matrigel plug angiogenesis assays showed significantly enhanced neovascularization in U19/Eaf2 knockout mice in response to VEGF (Fig. 6), providing evidence that U19/Eaf2 can modulate the vascular system presumably by binding to pVHL. At the same time, pVHL may also modulate the function of U19/Eaf2 because studies here show that the protein level of U19/Eaf2 is enhanced in the presence of VHL gene expression.

Like U19/Eaf2, pVHL has more than one binding partner; thus, U19/Eaf2- and pVHL-dependent signaling pathways are likely to be different but overlapping. A recent study identified pVHL as a binding partner of the tumor suppressor p53. Binding of pVHL led to an increase in both stability and transcriptional activation of p53, suggesting that other tumor suppressors might serve as mediators for the tumor-suppressive function of pVHL (39). In testes, the phenotypes are similar between the U19/Eaf2 knockout and VHL conditional knockout mice, suggesting significant functional overlaps between U19/Eaf2 and pVHL in this tissue. This also suggests that one possible mechanism for aspermogenesis in U19/Eaf2 knockout mice may involve pVHL. The function of pVHL in the testis is likely compromised in the absence of U19/Eaf2, which may cause a decrease in microtubule stability (40) and, subsequently, defects in both mitosis and meiosis leading to aspermogenesis (7). The functional overlap between the U19/Eaf2 and pVHL axes may be less extensive in other tissues, given that U19/Eaf2 knockout mice did not display similar phenotypes outside of the testes. Future studies will be required to elucidate the fundamental significance of the cross talk between U19/Eaf2 and pVHL in the context of other binding partners.

In summary, our novel findings strongly argue that U19/Eaf2 is a new regulator of the pVHL pathway and provide compelling evidence of cooperation between the two tumor suppressors, U19/Eaf2 and pVHL.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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Figure 6. The effect of U19/Eaf2 deletion on angiogenesis in Matrigel plugs s.c. implanted in mice. Left, representative digital images of microvessels, stained with anti-CD31 antibody, in Matrigel plugs in the presence of 2 ng/mL VEGF in the wild-type and U19/Eaf2 knockout mice. Right, quantitative analysis of the microvessel densities (MVD) in the Matrigel plugs.
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