Characterization of a von Hippel Lindau Pathway Involved in Extracellular Matrix Remodeling, Cell Invasion, and Angiogenesis

Ghada Kurban,1 Valérie Hudon,1 Eric Duplan,1 Michael Ohh,2 and Arnim Pause1

1McGill Cancer Center and Department of Biochemistry, McGill University, Montreal, Quebec and 2Department of Laboratory Medicine and Pathobiology, Faculty of Medicine, University of Toronto, Toronto, Ontario, Canada

Abstract

Inactivation of the von Hippel-Lindau (VHL) tumor suppressor gene results in highly vascularized tumors, making the VHL tumor syndrome an ideal system to study the mechanisms of angiogenesis. VHL operates along two pathways with the first involving hypoxia-inducible factor-α degradation and down-regulation of its proangiogenic target genes vascular endothelial growth factor and platelet-derived growth factor-β, and the second pathway promoting extracellular matrix (ECM) assembly. Secretion of proangiogenic factors was shown to be a primary inducer of angiogenesis. Here, we show that loss of ECM assembly correlates with tumor angiogenesis in VHL disease. Upon inactivation of the VHL-ECM assembly pathway, we observe tumors that are highly vascularized, have a disrupted ECM, and show increased matrix metalloproteinase-2 activity. Loss of the VHL pathway leading to hypoxia-inducible factor-α degradation results in tumors with increased vascular endothelial growth factor levels but with surprisingly low microvessel density, a tightly assembled ECM and low invasive ability. We conclude that loss of ECM integrity could promote and maintain tumor angiogenesis by providing a route for blood vessels to infiltrate tumors. (Cancer Res 2006; 66(3): 1313-9)

Introduction

von Hippel-Lindau (VHL) disease is an autosomal dominant familial cancer syndrome caused by the germ line mutation of the VHL tumor suppressor gene (1). Loss of VHL function leads to the development of highly angiogenic tumors including renal cell carcinoma (RCC), hemangioblastomas of the central nervous system, and pheochromocytoma (2–4). These highly vascularized tumors make VHL disease an ideal model to study angiogenesis.

Angiogenesis, which involves the growth of new capillaries from preexisting microvessels, is critical for various physiologic and pathologic processes, particularly tumorigenesis and metastasis. Ever since tumor growth was proposed to be dependent on angiogenesis (5), numerous studies have shown that it is required for the growth and metastasis of solid tumors that are over a few cubic millimeters in size (6). The angiogenic response, also known as the “angiogenic switch,” is initiated when the balance between the positive and the negative regulators of angiogenesis is disrupted in favor of the proangiogenic factors (6, 7).

The role of VHL in the regulation of proangiogenic factors stems from its ability to down-regulate the alpha subunit of the hypoxia-inducible factor (HIF-α), which exists in at least three isoforms (HIF-1α, HIF-2α, and HIF-3α; ref. 8). The VHL protein is part of an E3 ubiquitin ligase complex which targets HIF-α for ubiquitylation and degradation by the 26S proteasome under normal oxygen levels (1, 9–15). Under hypoxic conditions or on loss of VHL function, HIF-α is stabilized, resulting in transcription and accumulation of various HIF target genes such as vascular endothelial growth factor (VEGF), platelet-derived growth factor-β, transforming growth factor-α, and glucose transporter 1 (GLUT-1; refs. 1, 8). It was recently shown that HIF-2α is the isoform responsible for the induction of proangiogenic and growth factors (16).

In addition to HIF-α regulation, VHL also plays an important role in maintaining extracellular matrix (ECM) integrity (1, 17–19). VHL-negative cells lose the ability to assemble a fibronectin matrix whereas reintroduction of VHL restores assembly (17, 18). ECM assembly is an important regulatory element of blood vessel formation. Degradation of ECM by various proteases allows endothelial cell migration, leads to the release of ECM-sequestered angiogenic factors, and results in blood vessel infiltration (20, 21). VHL was reported to interact with a 200 kDa protein (9, 22) that was subsequently identified as fibronectin (17). This interaction is lost with all naturally occurring VHL mutants tested to date, stressing its importance in the VHL disease (1).

The significance of matrix disassembly in VHL disease is not well understood. However, it was reported that VHL neddylation is required for fibronectin matrix assembly and leads to suppression of tumorigenesis (23). Moreover, a recent study of VHL function in Caenorhabditis elegans confirmed that VHL operates along two independent pathways: one being dependent on HIF-α, and the other involving ECM regulation (24), suggesting evolutionary conservation of both pathways. Here, we used a cellular system in which the VHL-ECM assembly and VHL-HIF-2α degradation pathways are uncoupled to study their contribution to invasion, tumorigenesis, and angiogenesis.

We show that VHL leads to the assembly of a fibronectin and collagen type IV network which correlates with suppression of invasiveness in vitro as well as angiogenesis and tumorigenesis in vivo. VEGF expression in VHL tumors is not sufficient to promote and maintain angiogenesis unless it is coupled with an aberrantly remodeled ECM matrix suitable for endothelial cell migration and blood vessel infiltration. We conclude that loss of the VHL-ECM assembly pathway is likely to promote angiogenesis activation which could lead to highly vascularized VHL tumors. These results might have implications for the induction of angiogenesis in other carcinomas.

Materials and Methods

Cells and cell culture. VHL-negative human renal carcinoma cell line 786-O, as well as stable transfectants of 786-O, including pRC-HAVHL clones...
WT8 and WT7, pRC vector alone clone pRC3, pRC-HAVHL mutant L188V, pRC-HAVHL mutant R64P, and pRC-HAVHL pools of WT8 clones transduced either with an empty retroviral vector (WTE) or a retrovirus expressing a mutant P53IA HIF2α (WTPA) were a kind gift from Dr. William G. Kaelin (Dana-Farber Cancer Institute and Brigham and Women's Hospital, Harvard Medical School, Boston, MA). These cell lines have been described previously (18, 25). All the cells were grown in DMEM (Invitrogen, Burlington, Ontario, Canada) supplemented with 10% fetal bovine serum (FBS, Sigma-Aldrich Canada Ltd., Oakville, Ontario, Canada), 0.5 mg/ml G-418 in addition to 1.5 μg/ml of puromycin (in case of the retrovirally transduced cell lines) and maintained at 37°C in an atmosphere of 5% CO2.

**Antibodies.** The primary antibodies used were mouse anti-human fibronectin (BD Biosciences Pharmingen, Mississauga, Ontario, Canada), mouse anti-human collagen type IV (Chemicon, Temecula, CA), rabbit anti-human collagen type IV (Abcam, Cambridge, MA), rabbit anti-GLUT-1 (Alpha Diagnostic, San Antonio, TX), rabbit anti-HIF-2α (Novus, Littleton, CO), mouse anti-HA (12CA5; Roche, Laval, Quebec, Canada), rabbit anti-VEGF (Neomarkers, Labvision, Fremont, CA), and rat anti-mouse CD31 (BD Pharmingen). Mouse anti-MMP-2 was kindly provided by Dr. Rafael Friedman (Wayne State University, Detroit, MI). Secondary antibodies used were rhodamine-conjugated goat anti-mouse, FITC-conjugated goat anti-mouse, horseradish peroxidase–conjugated anti-mouse (Jackson ImmunoResearch Laboratories, Inc., West Grove, PA) as well as horseradish peroxidase–conjugated anti-rabbit (Amersham).

**Immunofluorescence microscopy.** Cells (5 × 10⁴) were grown on coverslips in six-well plates for 6 days. They were then washed with PBS, fixed with 95% ethanol at −20°C, followed by incubation with an antifibronectin monoclonal antibody. They were then washed with PBS-0.1% Tween and incubated with the secondary rhodamine-conjugated goat anti-mouse antibody. The nuclei were finally stained with 4′,6-diamidino-2-phenylindole (Sigma) and the cells were visualized using a Zeiss immunofluorescent microscope.

**Nude mouse xenograft assays.** To perform tumor inoculation, the different cell lines were harvested by trypsinization. Cell concentration and viability were determined by trypan blue staining, and 10⁷ cells resuspended in sterile PBS were injected s.c. into both flanks of 6- to 8-week-old CD1 nude (nu/nu) mice (Charles River Laboratories, Wilmington, MA). Nine to 10 weeks after injection, the animals were sacrificed and the s.c. tumors were excised. Bidimensional measurements were done using calipers and tumor volumes were calculated using the following formula: volume = width² × length / 2. The tumors were either embedded in optimal cutting temperature (Tissue Tek, Sakura Finetek USA, Inc., Torrance, CA) and stored at −80°C, or they were fixed in 4% paraformaldehyde/PBS overnight and embedded into paraffin for immunohistochemical analysis.

**Immunohistochemistry.** Ten-micrometer frozen sections of tumors were fixed in ice-cold acetone and rinsed in PBS. After blocking in PBS-0.05% Tween containing 1% (w/v) bovine serum albumin (BSA), sections were incubated with mouse anti-human collagen type IV and mouse anti-human fibronectin. The sections were then washed in PBS-0.05% Tween and incubated with FITC-conjugated goat anti-mouse IgGs. Finally, the sections were stained with 4′,6-diamidino-2-phenylindole hydrochloride (Sigma) and the cells were visualized using a Zeiss immunofluorescent microscope.

**Western blots.** For visualization of blood vessels, frozen sections were incubated with an antifibronectin antibody. For incubation with goat anti-rat secondary antibody (Vector Laboratories, Inc., Burlingame, CA), the signal was enhanced using an ABC kit (Vector) and color was developed with the chromophore diaminobenzidine (Sigma). Sections were counterstained with Fast Red nuclear (Sigma). Microvessel density was carried out from three different cell lines and was finally counterstained with hematoxylin.

**VEGF ELISA.** The various cell lines were seeded in six-well plates (5 × 10⁴ cells/well) and allowed to grow for 72 hours in normal growth medium. The conditioned medium was then removed, and VEGF levels were assayed by ELISA (R&D Systems, Minneapolis, MN) according to the manufacturer’s protocol. For normalization, the VEGF protein level was divided by the total cellular protein concentration for each sample.

**Figure 1.** Detection of fibronectin matrix assembly using indirect immunofluorescence. A, the various cell lines were grown on coverslips for 6 days. Fibronectin matrix assembly was verified by indirect immunofluorescence, using a monoclonal anti-fibronectin antibody. Cells expressing wild-type VHL (WT8, WT7, WTE, and WTPA) were able to assemble a distinguishable fibronectin matrix, in contrast to VHL-defective cells (786-0, pRC, L188V, and R64P). This phenomenon is regulated by VHL in a HIF-2α-dependent manner (magnification, ×40). B, HIF-2α expression. Using an anti-HIF-2α polyclonal antibody, HIF-2α levels were found to be elevated in cells lacking VHL (786-0 and pRC; lanes 1 and 2) as well as in WTPA cells which express a nondegradable form of HIF-2α (lane 6). This is in contrast to the low levels detected in WT8, WT7, and WTE cells (lanes 3, 4, and 5). Cells harboring type 2C VHL mutants, L188V and R64P, maintain the ability to degrade HIF-2α, resulting in barely detectable HIF-2α levels (lanes 7 and 8). Tubulin was used as a loading control. C, functionality of HIF-2α was assessed by verifying the expression levels of its target gene GLUT-1 using a polyclonal anti-GLUT-1 antibody. All cells overexpressing HIF-2α showed increased levels of GLUT-1. Actin was used as a loading control.
in terms of mean tumor volume, microvessel density, and VEGF protein observed by staining with 0.5% Coomassie blue.

The samples were normalized for intracellular protein concentration then collected and concentrated using Amicon centrifugal filters devices (Millipore, Nepean, Ontario, Canada). The samples were normalized for intracellular protein concentration then boiled in SDS-containing sample buffer. For the fibronectin and collagen IV secretion inhibitors. They were then boiled in SDS-containing sample buffer prior to loading on SDS-PAGE gels.

**Invasion and zymogram assays.** Polycarbonate inserts (8 μm pore, Fisher, Nepean, Ontario, Canada) were first coated with growth factor–reduced (GFR) Matrigel (BD Biosciences) diluted in serum-free DMEM. Cells (10^4) in complete medium (DMEM + 10% FBS) were then plated on the Matrigel layer and allowed to invade for 22 hours. They were then fixed with 10% neutral buffered formalin (Surgipath Canada, Inc., Winnipeg, Manitoba, Canada) and stained with 0.1% crystal violet solution (Sigma). The Matrigel layer was scraped off with a cotton swab followed by visualization using a light microscope. The experiments were repeated in triplicate and cell counts were based on four different fields per experiment.

Zymogram assays were done using conditioned medium from the various cell lines. Cells were grown in 0.1% BSA for 35 hours and the media was then collected and concentrated using Amicon centrifugal filters devices. The samples were normalized for intracellular protein concentration followed by PAGE on 0.1% gelatin gels. The gels were then washed in a solution of 2.5% Triton in 10 mmol/L Tris-HCl (pH 7.5) for 1 hour followed by a 15-minute wash with 10 mmol/L Tris-HCl (pH 8.0). They were then incubated in a zymogram containing 50 mmol/L Tris-HCl (pH 8.0), and 10 mmol/L CaCl$_2$ overnight at 37°C. Gelatinolytic activity was then observed by staining with 0.5% Coomassie blue.

**Statistical analysis.** Student’s *t* test was used to analyze the differences in terms of mean tumor volume, microvessel density, and VEGF protein levels. Statistical significance was accepted for a value of *P* < 0.05.

**Results**

**Derivatives of the 786-0 RCC cell line provide a system to study the VHL pathways involved in suppression of tumorigenesis and angiogenesis.** VHL operates along two pathways, the first leading to HIF-α degradation and the second resulting in ECM assembly. The role of VHL in suppressing tumor formation through HIF-2α regulation has been established, but it is not clear how loss of ECM assembly leads to tumor formation. We used the VHL-negative RCC cell line 786-0 and its derivatives to dissect and study these two pathways separately. 786-0 cells only express the HIF-2α isoform. We tested these RCC cell lines for their ability to assemble an ECM matrix and we observed that cells lacking VHL function failed to assemble a fibronectin matrix (Fig. 1A), whereas reintroduction of VHL restored fibronectin assembly as shown previously (17). To determine whether disruption of the VHL-HIF-2α degradation pathway abolishes fibronectin matrix assembly, we used WTPA cells which are VHL-positive (VHL+) cells retrovirally infected to produce a nondegradable form of HIF-2α. WTE cells, which are VHL+ cells infected with an empty virus, were used as control for WTPA cells (25). Our results showed that WTPA cells can assemble a fibronectin matrix similar to the VHL+ cells (Fig. 1A). Therefore, disruption of the VHL-HIF-2α degradation pathway does not affect the ability of RCC cells expressing WT-VHL to regulate ECM assembly. WTPA cells provide a system whereby only the VHL-HIF-2α degradation pathway is disrupted but the VHL-ECM assembly pathway is still functional. Cells expressing type 2C VHL mutants, L188V and R64P, failed to assemble a fibronectin matrix as described previously (18, 19). These VHL mutant–expressing cells provide an important tool to individually study the effect of loss of the VHL-ECM assembly pathway. The presence of HIF-2α was assessed in the different cell lines by Western blot (Fig. 1B) and its functionality was determined by the expression of one of its target genes, GLUT-1 (Fig. 1C).

**Inactivation of the VHL-ECM assembly pathway is involved in tumor formation.** It was reported that loss of ECM integrity influences cellular transformation and tumorigenesis in vivo (26). Therefore, we examined whether inactivation of the VHL-ECM assembly pathway contributes to tumorigenesis. To this end, we injected the various RCC cell lines s.c. into the flanks of nude mice. Nine weeks after the inoculation, tumor formation was observed with cells that have lost the VHL-HIF-2α degradation pathway (WTTPA) as well as with cells that have lost regulation of both VHL-ECM assembly and VHL-HIF-2α degradation pathways (pRC) as previously described (Fig. 2A; ref. 25). In contrast, cells expressing wild-type VHL (WT7) failed to form tumors. Interestingly, the cell lines defective in the VHL-ECM assembly pathway (L188V and R64P) also formed tumors that developed with a later onset than tumors derived from pRC and WTPA cells with the tumor incidence comparable for all cell types (Fig. 2A and B). These results suggest that VHL suppresses tumor formation by operating along both pathways: ECM assembly and HIF-2α degradation. Our data, using naturally occurring VHL mutants, substantiate the findings of Stickle et al. In their report, a VHL mutant defective for neddylation, which failed to assemble a fibronectin matrix but could still degrade HIF-α, lead to tumor formation (23).
Highly angiogenic tumors result from loss of the VHL-ECM assembly pathway and not the VHL-HIF-2α degradation pathway. VHL disease is characterized by the formation of highly vascularized tumors. In order to identify which pathway is responsible for angiogenesis in this RCC model, we examined blood vessel formation by staining the different tumors for an endothelial cell marker, CD31 (Fig. 3A). As previously observed, pRC cells, lacking both the VHL-HIF-2α degradation and VHL-ECM assembly pathways, formed highly vascularized tumors as compared with the ones formed by the cells lacking only the VHL-HIF-2α degradation pathway (WTPA; ref. 25). Very interestingly, tumors derived from cell lines lacking a functional VHL-ECM assembly pathway (L188V and R64P) were highly vascularized similar to tumors obtained with pRC cells (Fig. 3A). The differences in vascularization were confirmed by quantification of the microvessel density for each type of tumor (Fig. 3B). A three times more significant decrease was observed in the microvessel density of the WTPA tumors as compared with the other types of tumors (P < 0.01 versus pRC; P < 0.001 versus L188V and R64P). Poor vascularization and low microvessel density comparable to the WTPA tumors were also observed in other RCC tumors derived from cells expressing wild-type VHL, SKRC-39 (data not shown). Necrotic areas which were mainly detected in WTPA tumors were probably due to lack of vascularization (Fig. 3A, arrow). These results suggest that the development of a vascular network following pVHL inactivation occurs upon loss of the VHL-ECM assembly pathway and is not only dependent on the VHL-HIF-2α degradation pathway.

Expression and secretion of VEGF is not sufficient to develop a vascular network in RCC tumors. Tumors secrete a variety of angiogenic factors including VEGF. To examine if the high level of angiogenesis observed with the VHL tumors was due to an increase in VEGF levels, we first measured secreted VEGF protein levels in the different cell lines using an ELISA assay. Due to the induction of VEGF expression by increased HIF-2α levels, WTPA, as well as pRC cell lines had a significant 2-fold increase in secreted VEGF levels over cell lines expressing wild-type VHL (WT7) and those that have lost the VHL-ECM assembly pathway only (L188V and R64P; Fig. 4A). The higher VEGF levels in the pRC and WTPA cells were due to induction of VEGF expression by increased HIF-2α levels. We then did indirect immunostaining on paraffin sections of the different tumors to verify the VEGF status in vivo and observed similar VEGF staining in all tumors (Fig. 4B).

Figure 3. Determination of the VHL pathway involved in angiogenesis. A, representative histologic sections stained with H&E for common tumor histology. Note that WTPA tumors present areas of necrosis (arrow) in contrast to the tumors defective for the VHL-ECM assembly pathway. To analyze the vascularization of the various tumors, frozen sections were stained with an antibody against CD31, pRC cells, defective in both the VHL-HIF-2α degradation and VHL-ECM assembly pathways, as well as L188V and R64P cells, which are defective only in the VHL-ECM assembly pathway showed increased CD31 staining as compared with WTPA tumors defective in the VHL-HIF-2α degradation pathway only. B, quantification of CD31 levels was done blindly and is presented in terms of area fraction occupied by the microvessels per microscope field (expressed as a percentage). pRC tumors (4.5 ± 1.7%), defective in the HIF-2α pathway, displayed significantly fewer microvessels as compared with the other three cell types, pRC (13.9 ± 4.8%), L188V (15.5 ± 8.6%), and R64P (11.5 ± 4.7%). **, P < 0.01 versus pRC group (magnification, x20).

Figure 4. Expression of VEGF protein in cells and tumors. A, 5 × 10⁶ cells of the various cell lines were plated on six-well plates and allowed to grow in normal growth medium. Conditioned supernatant was harvested after 72 hours, and VEGF protein levels analyzed by an ELISA assay. VEGF values were normalized to total cellular protein concentration and represent the mean from two independent experiments. VEGF protein levels are elevated in pRC (12.7 ± 1.6) and WTPA (12.1 ± 1.6) cell lines as compared with WT7 (5.9 ± 0.3), L188V (4.8 ± 0.6) and R64P (5.5 ± 0.6). ***, P < 0.001 versus pRC group. B, histochemical analysis of VEGF expression in the various tumors. All tumors showed comparable VEGF staining (magnification, x100).
Tumors formed upon loss of the VHL-ECM assembly pathway display a loosely remodeled ECM. Because VEGF is expressed in all tumors, it was surprising that WTPA tumors were so poorly vascularized as compared with L188V and R64P tumors. It was shown that VEGF can be sequestered by ECM proteins and released upon ECM remodeling by matrix metalloproteinases (MMP; refs. 21, 27). Therefore, we examined ECM deposition within tumors formed by the different cell lines. Because fibronectin and collagen type IV are two major proteins of the ECM, we did immunofluorescence staining of both proteins on frozen sections of the various tumors (Fig. 5A). The fibronectin and collagen type IV matrices within the tumors derived from cell lines that have lost the VHL-ECM assembly pathway (L188V and R64P) and those that have lost both VHL-HIF-2α degradation and VHL-ECM assembly pathways (pRC) were loose and aberrant as opposed to tumors formed by WTPA cells that have lost the HIF-2α degradation pathway (Fig. 5A). The latter tumors exhibited a dense and organized fibronectin and collagen type IV network that was evenly distributed throughout the tumor sections. This ECM organization is similar to other RCC tumors derived from the SKRC-39 cell line (wild-type VHL; data not shown). These results suggest that dysregulation of the VHL-ECM assembly pathway, leading to an aberrant ECM, is required for the distribution of blood vessels throughout VHL tumors.

Disorganization of the ECM in VHL tumors does not result from a defect in expression or secretion of ECM proteins. We tested whether there was a defect in expression or secretion of fibronectin and collagen type IV which would explain why VHL mutant cells fail to assemble an ECM. We did Western blot analysis using total cell lysates from the different cell lines. All the cell lines tested, including VHL mutant cell lines, produced fibronectin and collagen type IV (Fig. 5B). The expression levels of fibronectin and collagen type IV varied among the different cell types and showed no correlation with the ability of cells to assemble an ECM. To verify whether there was a defect in secretion of the two proteins in the cells expressing mutant forms of VHL, we did a Western blot analysis of secreted fibronectin and collagen type IV in the cell culture media. Although they were unable to assemble an ECM, VHL-defective cells, similar to cells expressing wild-type VHL, maintained the ability to secrete fibronectin and collagen type IV (Fig. 5C).

Loss of the VHL-ECM assembly pathway results in an invasive behavior of cells. Tumor angiogenesis requires ECM degradation which enables blood vessels to penetrate the basal lamina and invade the tumor tissue. It was shown that cells lacking VHL are invasive whereas reintroduction of VHL suppresses their invasive ability (28). We tested the ability of the different VHL mutant cell lines to invade GFR Matrigel. Our results showed that cells that have lost the VHL-ECM assembly pathway (L188V and R64P), and cells that have lost both the VHL-ECM assembly and VHL-HIF-2α degradation pathways (786-0 and pRC) were highly invasive (Fig. 6A) and expressed high MMP-2 levels as determined by Western blot analysis (Fig. 6B). The MMP-2 protein levels correlated with MMP-2 activity when tested in zymogram assays (Fig. 6C). Conditioned medium from primary rat smooth muscle cells, which express high levels of MMP-2, were used as a positive control (data not shown). Reintroduction of wild-type VHL in WT8, WT7, and WTE cells suppressed their invasive capacity by at least 4-fold. Cells that have only lost the VHL-HIF-2α degradation pathway (WTPA

Figure 5. Analysis of ECM proteins in tumors and cells. A, tumor sections were stained for the ECM proteins, fibronectin and collagen type IV. WTPA cells, defective for the VHL-HIF-2α degradation pathway, displayed a dense and tightly organized network of fibronectin and collagen type IV. In contrast, pRC cells, lacking both the VHL-HIF-2α degradation and VHL-ECM assembly pathways, as well as L188V and R64P cells, which are defective only in the VHL-ECM assembly pathway, displayed a loose network of both ECM proteins (magnification, ×20). B, fibronectin and collagen IV expression. Cells were lysed using EBC buffer and total lysates were loaded on SDS-6% PAGE gels. All cells expressed significant levels of fibronectin and collagen IV regardless of their VHL status. Actin and tubulin were used as loading controls. C, fibronectin and collagen type IV secretion. Cells were grown in DMEM 0.1% BSA for 35 hours, the medium was then collected and concentrated, and then loaded on SDS-6% PAGE gels. All cells secreted fibronectin and collagen IV with variable patterns regardless of their VHL status.
cells) were significantly less invasive with diminished MMP-2 expression and activity, suggesting that loss of the HIF-2α degradation pathway does not promote invasiveness in this RCC setting (Fig. 6A and B). These data suggest that loss of the VHL-ECM assembly pathway leads to an increase in cell invasion and ECM degradation resulting in an aberrant tumor matrix and promotion of tumor angiogenesis.

Discussion

VHL inactivation leads to the development of tumors densely infiltrated with blood vessels. VHL functions along two pathways: the first leading to tumor suppression by HIF-2α degradation, and the second involving ECM regulation. HIF-2α controls many cellular processes including stimulation of VEGF expression which leads to increased angiogenesis (16, 29, 30), and induction of serum-independent cell proliferation (31). The high level of vascularization of VHL tumors has been attributed to VEGF overexpression. By binding to its receptors, VEGF could trigger an angiogenic response by acting on the proliferation, migration, and survival of endothelial cells (32). However, it has been reported that tumor vascularization upon VHL loss is a result of more than just HIF-α up-regulation (25). Other reports showed that the loss of HIF-1α negatively affects tumorigenesis (16, 33) and reduction in tumor growth was not due to a decrease in vascularization (33).

Because the function of the VHL-ECM assembly pathway is poorly understood, we verified its role in suppression of tumorigenesis and angiogenesis using a cell system where the two pathways (VHL-HIF-2α degradation and VHL-ECM assembly) were uncoupled. Loss of the VHL-ECM assembly pathway led to the formation of tumors. This is in contrast to a study by Hoffman et al. (18) where L188V cells did not form tumors. This discrepancy could be due to a shorter incubation time in nude mice. However, our data are in accordance with Stickle et al., who used a VHL mutant defective for fibronectin matrix assembly (23). In addition, tumors observed upon loss of the VHL-ECM assembly pathway (L188V and R64P) were highly angiogenic and such important vascularization was not observed in tumors overexpressing HIF-2α, although they secreted 2-fold more VEGF in vitro. These latter tumors grew quickly, suggesting that loss of the VHL-HIF-2α degradation pathway is required for mediating the growth of VHL tumors probably due to overexpression of transforming growth factor-α and cyclin D1 (16). On the other hand, loss of the VHL-ECM assembly pathway seems to be required for full-scale tumor angiogenesis in this RCC setting.

Loss of the VHL-HIF-2α degradation pathway does not disrupt the fibronectin and collagen type IV network whereas loss of the VHL-ECM assembly pathway leads to an aberrant ECM assembly and stimulation of angiogenesis. Remodeling of the ECM during tumor angiogenesis is triggered by secreted proteases such as MMPs (34) and these were found to play an important role in angiogenesis. MMPs could increase the release and activation of matrix-sequestered angiogenic factors, such as VEGF (21, 27). VEGF has been documented to be an important inducer of tumor angiogenesis and a 2-fold increase in its expression level was shown to cause an angiogenic response using an in vitro angiogenesis assay (20). Therefore, it is possible that in VHL tumors lacking the VHL-HIF-2α degradation pathway, the intact ECM serves to sequester the secreted VEGF thus hindering it from stimulating proliferation and migration of endothelial cells (WTPA tumors). Remodeling of the ECM by MMP-2, which is activated upon loss of the VHL-ECM assembly pathway, might result in the release of VEGF to exert its angiogenic effect.

As a result of their remodeling activities, MMPs were shown to expose cryptic sites within collagen type IV and this is associated with a gain of binding to αvβ3 integrin on endothelial cells and stimulation of angiogenesis (35, 36). Disruption of MMP-2 binding to αvβ3 was shown to inhibit tumor growth and vascularization in vivo (37). Moreover, it was shown that angiogenesis and tumor growth are suppressed in MMP-2 knockout mice (38). Therefore, it is possible that in tumors which have lost the VHL-ECM assembly pathway, remodeling of the ECM by MMP-2 allows exposure of cryptic sites within ECM proteins which could stimulate tumor growth and angiogenesis.

In conclusion, we propose that loss of the VHL-ECM assembly pathway could have several consequences: (a) removal of a barrier to facilitate growth and invasion of blood vessels, (b) release of ECM sequestered factors such as VEGF, (c) presentation of cryptic sites on ECM proteins resulting in increased neoangiogenesis. Loss of VHL function drives tumorigenesis and angiogenesis along,
at least, the VHL-HIF-2α degradation and VHL-ECM assembly pathways. Although HIF-2α activation leads to VEGF overexpression, our results suggest that progression of the complete tumor angiogenic response is observed on loss of the VHL-ECM assembly pathway resulting in extensive VHL tumor vascularization. These results stress the importance of elucidating the mechanism of the VHL-ECM pathway which seems to be involved in tumorigenesis, angiogenesis, and cell invasion.

Acknowledgments

Received 7/20/2005; revised 11/7/2005; accepted 11/23/2005.

References

Characterization of a von Hippel Lindau Pathway Involved in Extracellular Matrix Remodeling, Cell Invasion, and Angiogenesis

Ghada Kurban, Valérie Hudon, Eric Duplan, et al.


Updated version
Access the most recent version of this article at:
http://cancerres.aacrjournals.org/content/66/3/1313

Cited articles
This article cites 38 articles, 17 of which you can access for free at:
http://cancerres.aacrjournals.org/content/66/3/1313.full.html#ref-list-1

Citing articles
This article has been cited by 24 HighWire-hosted articles. Access the articles at:
/content/66/3/1313.full.html#related-urls

E-mail alerts
Sign up to receive free email-alerts related to this article or journal.

Reprints and Subscriptions
To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.

Permissions
To request permission to re-use all or part of this article, contact the AACR Publications Department at permissions@aacr.org.