Decrease in VEGF Expression Induces Intussusceptive Vascular Pruning

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Objective—The concept of vascular pruning, the “cutting-off” of vessels, is gaining importance due to expansion of angio-modulating therapies. The proangiogenic effects of vascular endothelial growth factor (VEGF) are broadly described, but the mechanisms of structural alterations by its downregulation are not known.

Methods and Results—VEGF_{165}+releasing hydrogels were applied onto the chick chorioallantoic membrane on embryonic day 10. The hydrogels, designed to completely degrade within 2 days, caused high-level VEGF presentation followed by abrupt VEGF withdrawal. Application of VEGF resulted in a pronounced angiogenic response within 24 hours. The drastic decrease in level of exogenous VEGF-A within 48 hours was corroborated by enzyme-linked immunosorbent assay. Following this VEGF withdrawal we observed vasculature adaptation by means of intussusception, including intussusceptive vascular pruning. As revealed on vascular casts and serial semithin sections, intussusceptive vascular pruning occurred by emergence of multiple eccentric pillars at bifurcations. Time-lapse in vivo microscopy has confirmed the de novo occurrence of transluminal pillars and their capability to induce pruning. Quantitative evaluation corroborated an extensive activation of intussusception associated with VEGF withdrawal.

Conclusion—Diminution of VEGF level induces vascular tree regression by intussusceptive vascular pruning. This observation may allude to the mechanism underlying the “normalization” of tumor vasculature if treated with antiangiogenic drugs. The mechanism described here gives new insights into the understanding of the processes of vasculature regression and hence provides new and potentially viable targets for antiangiogenic and/or angio-modulating therapies during various pathological processes. (Arterioscler Thromb Vasc Biol. 2011;31:2836-2844.)

Key Words: angiogenesis ■ morphogenesis ■ vascular biology ■ intussusceptive vascular pruning

The inaugural capillary plexus of any vascular network undergoes substantial remodeling for the vasculature to meet the needs of the developing organ and to establish the organ-specific angio-architecture.\(^1,2\) Vascular pruning is an essential adaptive mechanism resulting in regression of excessive vascular branches and creation of hierarchical, thermodynamically efficient angioarchitecture. This process has been known for many decades; in 1873 Roguet nicely illustrated this important phenomenon and named it “retraction of capillaries.”\(^3\) In 1961 Ashton stated that retraction of capillaries plays a key role in the reorganization of the primitive immature capillary network.\(^4\) It was considered that some channels become redundant, blood flow within them ceases, the lumen obliterates, and the endothelial cytoplasm and nuclei retract into the parent capillaries leaving behind a trail of cytoplasm that disappears.\(^5\) This was the first profound description of a mechanism of vascular pruning at capillary level. In the recent past, the understanding of vascular remodeling has been substantially improved, partially, through detailed investigations and descriptions of the process of intussusceptive angiogenesis, and in particular, intussusceptive vascular pruning (IVP).\(^1,6,7\) Our previous morphological studies have revealed the vasculature of different organs to undergo 2 main phases of development: an early sprouting phase followed by an intussusceptive phase, during which capillary sprouting is superseded by transcapillary pillar formation. Intussusception results in rapid expansion of the capillary network and the genesis of an organ-specific vascular tree, as well as its dynamic adaptation and intussusceptive branching remodeling (see review\(^1\)). An important facet of intussusception is the IVP. It can be summarized as the formation of multiple eccentric pillars at the bifurcation points and their subsequent successive fusions, which leads to partial and later to total luminal obstruction and separation (cutting-off) of one of the affected daughter branches. The

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latter mechanism is alternative to the one described by Ashton and, in contrast, it is not limited to the capillary level but also involves larger vessels. Vascular pruning results in reduction of the number of vascular branches and vascular density respectively. This is one of the mechanisms allowing the vascular system to dynamically adapt to the changing hemodynamical and metabolic influences and to create mature hierarchical and even more efficient angioarchitecture.8

The remodeling process is regulated by mechanical influences (hemodynamics), as well as by numerous not yet well-defined morphogens and cytokines. The most potent and best known angiogenic factor, vascular endothelial growth factor (VEGF)-A, plays a major regulatory role in the formation of blood vessels and is known to control proliferation and migration of endothelial cells.5–10 Proliferation of endothelial cells is regulated by absolute concentration of VEGF, whereas their migration and differentiation toward different functionalities depends on the concentration gradient of different VEGF-isoforms.11,12 VEGF also influences the survival of vascular endothelial cells.9 Our ongoing studies in the developing kidney revealed the relation between VEGF downregulation and intussusceptive pruning (A. Makanya et al, unpublished data). The sustained VEGF expression or, respectively, its inhibition was shown to play an important role in vascular remodeling.13,14 The finding by Kesht’s group that VEGF is the molecular link between hypoxia and angiogenesis was a milestone in the field of hypoxia-driven angiogenesis.15 In a previous study we demonstrated that inhibition of VEGF/VEGF-R signaling pathway led to vessel degeneration in the chick chorioallantoic membrane (CAM), most probably by means of vascular pruning.16 Moreover, in a recent study we described how application of a VEGF inhibitor in the mouse xenografts tumor model led to a temporary switch in the angiogenesis mode: from sprouting to intussusception.17 The described findings underpin the paramount role of VEGF (or actually its substrate sequences for FXIIIa as previously described.17 Hydrogels without addition of VEGF-A165 were used as vehicle controls. In cases where the applied gels contained FITC labeled peptides, we injected up to 0.1 mL of 3% Rhodamine B isothiocyanate dextran solution [molecular weight: 70’000 Da (Sigma, Buchs, Switzerland)] in isotonc NaCl solution.

**Materials and Methods**

**Local Application of VEGF-A**

A 1.0 μg recombinant VEGF-A165 protein (Peprotech, London) was administered in a 10 μL 2% (weight /Volume) synthetic fibrin analog hydrogel. The hydrogels were formed under physiological conditions by enzymatic cross-linking of 8-arm star-shaped poly (ethylene glycol) PEG polymers, which are functionalized with substrate sequences for FXIIIa as previously described.17 Hydrogels without addition of VEGF-A165 were used as vehicle controls. In order to facilitate the localization of the applied hydrogel and estimate the level of its degradation on the CAM in some experiments FITC-labeled peptides Lys-FITC (Ac-FKGGK-fluorescein-NH2, NeoMPS, Strasbourg, France), which are covalently tethered to the forming hydrogels, were added.

**Intravitral Microscopy of CAM Vasculature**

Chorioallantoic membranes were obtained by incubating chick embryos using the established shell-free culture method.18 On E10 portions of the CAM surface were treated with hydrogels that either contained or did not contain 1 μg of VEGF-A165. Twenty four, 48, and 72 hours after the application, the CAMs were intravenously injected with up to 0.1 mL of 3% FITC-dextran solution [molecular weight: 2’000’000 Da (Sigma, Buchs, Switzerland)], prior to inspection in an epifluorescence microscope (Polyvar-Reichert, Glattbrugg, Switzerland) equipped with a Canon 5D Mark II camera for both video recording and taking of still images. The still images and video sequences of at least 4 fields of view were taken per application site for further quantitative evaluation. The total number of branching points per field of view was assessed using analySIS Software 5.0 (Soft Imaging System, Muenster, Germany) by means of user-driven skeletonization of the vasculature.

**Determination of Remaining VEGF by Enzyme-Linked Immunosorbent Assay**

We harvested and homogenized 1 cm² sized CAM samples from the site of treatment (including remaining hydrogel mass) in 500 μL lysis buffer (PBS pH 7.2, 0.1% Tween, 0.2% BSA, 1 mmol/L Benzamidine, 20 mmol/L NaF, 1 mmol/L PMSF, 2% Protease Inhibitor Cocktail Set III [Calbiochem, Darmstadt, Germany]) and incubated for 24 hours at 4°C on a rotating wheel. VEGF-A165 content in the lysis buffer was measured using a human VEGF enzyme-linked immunosorbent assay (ELISA; PeproTech, London, UK) according to the manufacturer’s protocol.

**Semithin Serial Sectioning**

CAM samples of sites of interest were harvested and fixed in 2.5% (v/v) glutaraldehyde solution buffered with 0.03 mol/L potassium phosphate (pH 7.4, 370 mOsm). They were then postfixed in 1% OsO4 [buffered with 0.1 mol/L sodium cacodylate (pH 7.4, 340 mOsm)], dehydrated in ethanol, and embedded in epoxy resin. Thousands of 0.8-μm-thick serial sections were prepared using glass knives and stained with Toluidine Blue. The serial sections were viewed and images captured at low magnification using a light microscope (Leica, Letz DM), equipped with Leica DFC480 camera. The set of images obtained was aligned using Adobe Photoshop CS3 Software and imported as stacks into Imaris Software for low resolution in an epifluorescence microscope (Polyvar-Reichert, Glattbrugg, Switzerland) by means of user-driven skeletonization of the vasculature. The Multiple Image Alignment tool of the aforementioned software was used to obtain overview images representing the whole application site with degrading gel at different timepoints. In cases where the applied gels contained FITC labeled peptides, we injected up to 0.1 mL of 3% Rhodamine B isothiocyanate dextran solution [molecular weight: 70’000 Da (Sigma, Buchs, Switzerland)] in isotonc NaCl solution.

**Vascular Casting**

Vascular casts were prepared as previously described.19 Briefly, CAM vasculature was perfused with a freshly prepared solution of Mercrox® (Vilene Company, Japan) containing 0.1 mL of accelerator per 5 mL of resin. One hour after perfusion, the CAM were transferred to 7.5% potassium hydroxide for dissolution of tissue, which was effected over a course of 2 to 3 weeks. After washing, the casts were dehydrated in ethanol and dried in a vacuum desiccator. The samples were then sputtered with gold to a thickness of 10 nm and examined in a Philips XL-30 SFEG scanning electron microscope.

**Index of Intussusception and Statistical Analysis**

From video sequences the number of branching points involved in intussusception (ie, those showing pillars) could be assessed. The
vasculature skeletons on the corresponding still images were used to find out the total number of branching points per each investigated field of view. By dividing the number of intussusception involved branching points by the number of branching points, the index of intussusception could be determined. At least 4 fields of vision per application site were evaluated (n/11005 4 per time point and per group, at least 16 fields of vision per time point and group) The used magnification was /11003 40. Statistical analysis was done using the 2-tailed t test. The difference was considered significant if P/11021 0.05.

Apoptosis Assay
Apoptotic cells were detected in vivo using Annexin V-FITC apoptosis detection kit (Sigma-Aldrich Chemie GmbH, Steinheim, Germany). The solution was injected directly into the CAM vascular system. After 15 minutes of incubation the vasculature was examined using a Leica AF 6000LX fluorescence microscope equipped with the Leica DFC 360FX digital camera. Annexin V-positive (=apoptotic) cells appeared as green spots in the images obtained.

Results

VEGF Release Profile
An angiogenic response was stimulated by local application of gels loaded with 1 μg of VEGF or empty gels (control) on top of growing chick CAM at E10. To allow the estimation the VEGF release profile we have added Lys-FITC to the gel composition. The FITC labeling of the hydrogel enabled the visual presentation of the proteolytic gel degradation and allowed the correlation of proteolysis-triggered release of VEGF with the amount of the remaining hydrogel mass20 (Figure 1). Another advantage of this application method is that removal of the gel after certain application time was not necessary in our experiments, avoiding the very often observed nonspecific reactions of the CAM vasculature due to mechanical damage or irritation.21 The gel was considerably degraded as early as 24 hours and completely degraded in 48

![Figure 1](image-url)
hours after the application (Figure 1A–1C) and no more detectable at later time-points (data now shown).

Determination of the VEGF-A levels using ELISA-kit has corroborated that the amount of exogenous VEGF-A remained at a significantly higher level at 24 hours when compared to controls. At 48 hours VEGF expression decreased back to the level in control samples (Figure 1D).

Thus, in our model an exhaustive stimulation by exogenous VEGF was achieved for more than 24 hours followed by a drastic drop in VEGF levels by 48 hours.

**Morphogenesis of IVP**

In order to investigate the detailed morphology of the affected vasculature we made vascular corrosion casts of CAMs at 24 and 48 hours after VEGF application. Observation of casts using scanning electron microscope corroborated the hypothesis that besides the differences between control (Figure 2a) and VEGF-treated samples, there was a qualitative change between the vasculature during VEGF stimulation (at 24 hours) (Figure 2B) and when the VEGF presentation was discontinued, namely at 48 hours (Figure 2C–2H). In the latter ones there were numerous bifurcations involved in IVP. Various stages of the pruning process were documented on vascular casts made at 48 hours after VEGF-A application (Figure 2C–2H). Initially single eccentrically positioned pillars (arrowheads in Figure 2) arise (Figure 2C–2D), they then grow in number and girth (Figure 2D), and folds (arrows in Figure 2) emerge between them (Figure 2E–2F). As the folds become deeper (Figure 2F–2G), pillars fuse and finally lead to the “cutting-off” of the corresponding vessel branches (Figure 2G). These changes take place simultaneously in a “cascade-like” manner at many branching levels, from capillary plexuses up to larger vessels (Figure 2H). Note the IVP-altered capillary plexus indicated with an asterisk in Figure 2H.

Apparently, IVP was activated 48 hours after VEGF-A application, soon after the dramatic decrease in VEGF-A expression.

**Time Course of IVP**

Time-lapse in vivo video microscopy (Figure 3) confirmed the aforementioned activation of IVP following VEGF-A downregulation by revealing the appearance of many asymmetrically positioned pillars, the hallmarks of intussusceptive pruning. The fast appearance and progression of the pillars (within hours) enabled the documentation of various stages of IVP in vivo: from appearance of single eccentrically positioned pillars to formation of pillar rows (Figure 3A–3C) and almost complete “cutting-off” of the daughter vessel with drastic decrease in the size of its orifice resulting in the vessel being hardly patent. The same fate befell the neighboring capillary plexus (Figure 3D–3F). The corresponding vascular cast image with multiple intussusception involved branching and, due to pruning altered capillary plexus, is presented in Figure 2H.

These findings illustrate the rapid change in vascular structure accomplished by intussusception.

**High Resolution 3D-Reconstruction Using Serial Semithin Sectioning**

The branching point of interest with emerging pillars observed in vivo (Figure 4A–4C) was fixed just after the last in vivo observation and then processed for semithin sectioning. Over 2000 serial 0.8 μm-thick sections were made with the cutting plane being perpendicular to the CAM surface and stained with toluidine blue. Low magnification images of these sections were acquired, aligned, and processed for low resolution 3D-reconstruction using Imaris Software. The data obtained by 3D-reconstruction was used for precise localization of the vascular segment of interest (white rectangle in Figure 4C) by superimposing it with the in vivo images. Afterward high magnification images of determined sites in 140 sections were acquired for high resolution 3D-
reconstruction: the virtual section through the 3D-stack is presented in Figure 4D. Further, the 3D surface of the vascular lumen of the in vivo segment of interest was computer generated (Figure 4e) and corresponded to the in vivo obtained images (see Figure 4C for comparison). The intraluminal view demonstrates clearly that the structures observed in vivo were emerging pillars (Figure 4F). On the corresponding semithin sections, the intraluminal protrusions of endothelial cells at the opposing sides of the vascular lumen were apparent (Figure 4G–4H).

**Adaptation (Regression) of the Vasculature Happens at Different Hierarchical Levels, and Including Larger Supplying Vessels**

At 72 hours after VEGF application even the usually rigid larger arteries also underwent the intensive process of adaptation (Figure 5). Intussusceptive pruning and remodeling was not restricted to distal part of vasculature but could be observed along the entire vascular tree (Figure 5B–5D). This presents the integrity of the vasculature system adaptation and involvement of intussusception in the regressive remodeling at all hierarchical levels of the vasculature. The intraluminal protrusions of endothelial cells at the opposing sides of the vascular lumen were apparent (Figure 4G–4H).

**Morphometry of the Affected Vasculature**

Quantification of intussusception was done using the data obtained using in vivo microscopy. The parameter we have chosen as the measure of intussusception was the index of intussusception involved branching points. The results obtained have corroborated the qualitative description: In the control areas, there were no significant changes of this parameter at all 3 time-points. On contrary, the VEGF-A treated areas were characterized by a constant increase of intussusception involved branching points with the most prominent value (>2.5fold in comparison with control) at 72 hours (Figure 6).

**Apoptosis of Endothelial Cells at Remodeled Sites**

To address the question whether the endothelial cells undergo apoptosis we made the additional experimentation: The IV injection of FITC-labeled anti-Annexin–V apoptosis detection kit was made prior in vivo observation of the VEGF application sites. The apoptotic cells could be found at 48 hours and more after the application. At E10+48 hours there were scattered apoptotic cells in the capillary plexus and at E10+72 hours one could observe already bigger vessels with numerous Annexin-V–positive cells (see Figure 7).

**Discussion**

**Mechanism of Vascular Pruning**

The mechanism of IVP has been rather explicitly described in the last decade. At the initial stage intraluminal protrusions of endothelial cells lying at the opposing sides of the vessel are formed. The contact between them has to be established for the single pillar to emerge. The peculiarity of IVP is that the pillars are positioned eccentrically from the axis of the mother vessel. During the progression of IVP the pillars increase in number and size, they fuse by folds, and finally lead to complete “cutting-off” of one of the daughter vessels. Importantly, this process that leads to regression of the vasculature takes place at all levels of the vascular bed, from capillary plexus to larger vessels (see Figure 2 and Figure 5).
VEGF and IVP

**VEGF Application Stimulates Angiogenesis in the CAM**
In the present study we could observe typical signs of stimulated angiogenesis following application of VEGF-A. It is well described that in addition to the amount of the applied or expressed VEGF, its release modality, i.e., slow versus rapid liberation, is a crucial factor for the angiogenic response and vascular morphogenesis.\textsuperscript{23–26} Considering the data in the literature, the fibrin gel application was chosen to create an interface between CAM and VEGF-A reservoir and allow a sustained release during the proteolytic degradation of the gel. Microscopical and ELISA-based analysis suggest an exogenous VEGF presentation that lasts for approximately 24 hours after application and that once the gels are completely degraded rapidly terminates. The reversion to the physiological level at 48 hours (on E12) is confirmed by VEGF-ELISA.

**VEGF-A Withdrawal Leads to IVP**
Although decrease of VEGF concentration has been previously presumed to induce vascular remodeling and pruning,\textsuperscript{6,27,28} this hypothesis could only now be affirmed in the present study both qualitatively and quantitatively. Within a rather short time after the decrease of VEGF concentration we observed appearance of many asymmetrically positioned pillars at branching points. Our observations indicate that intussusceptive pillar formation is a major mechanism in VEGF-dependent vascular pruning. This finding is in accordance with other studies describing induction of intussusception and degeneration of immature vessels after inhibition of VEGF-signaling\textsuperscript{14,16,29} or hyperoxia-induced withdrawal of VEGF.\textsuperscript{30,31} In contrast to the mentioned studies we show here that (VEGF-dependent) IVP is not limited to the immature vasculature but takes places at higher hierarchical levels of the mature vascular bed (see branching of the larger arteries).
in Figure 5). We believe that the IVP of larger mature vessels may be secondary to hemodynamic changes in distal vascular segments, ie, capillary plexus. There are controversial data in the literature about the fate of the occluded vascular segments. Although most authors postulate that the process involves apoptosis, pruned ECs may also reassemble into other vessels or dedifferentiate. In our model the process of vascular pruning definitely involves apoptosis (Figure 7) but involvement of other mechanisms cannot be excluded.

Clinical Implications
Inhibition of angiogenesis is a promising strategy for treatment of cancer and other disorders, whereas promoting growth of new vessels is important for treatment of ischemic disorders. Most antiangiogenic agents in tumor treatment are targeting VEGF signal transduction. The antiangiogenic treatment was supposed to deprive the supply of oxygen and nutrients to the tumor by destroying its vasculature. Even a single infusion of bevacizumab, a VEGF-specific antibody, was shown to significantly reduce the microvascular density in rectal carcinoma patients. However, the survival benefits of antiangiogenic drugs have thus far been rather disappointing, stimulating interest in developing more effective ways to combine antiangiogenic drugs with established chemotherapies. In the recent past it has been proved that an antiangiogenic approach, when combined with chemotherapy, results in increased survival in patients with advanced malignancies. The improved efficacy is nowadays often explained by “vascular normalization.” The hypothesis of transient “normalization” of the vasculature states that some antiangiogenic drugs improve tumor perfusion by reducing

![Figure 5. Micrographs from in vivo imaging showing the cascade-like intussusceptive remodeling of the arterial tree after VEGF down-regulation. A low magnification image of an artery (Ar) with several branchings is shown in A. The branchings undergoing intussusceptive remodeling and pruning (white rectangles in A) are displayed at higher magnification in panels B–D. The pillars and meshes evident on the vessels (arrowheads in B–D) lead to simultaneous IVP and remodeling (dashed lines in B–D) at more proximal and distal branching points. This shows the integrity of vascular tree adaptation by means of intussusceptive pruning following vascular endothelial growth factor (VEGF)-A downregulation. The branches separated via IVP are indicated by asterisks in B–D. This figure supports the data shown in Fig. 2H that remodeling takes place at different hierarchical levels and is not limited to venous compartment.](image)

![Figure 6. Quantitative evaluation of intussusceptive pruning at branching points. The graph represents the index of intussusception at 24, 48, and 72 hours after vascular endothelial growth factor (VEGF) gel application. The VEGF-A treated areas were characterized by an outstanding increase (P<0.01) of the intussusception involvement at 72 hours, at the time-point when VEGF-A expression was already significantly decreased. In the controls there was no significant change between the time-points. The values represented in the graph are mean±SD (n=4).](image)
vascular density, stabilizing the vessel wall (perivascular cell and basement membrane coverage), reducing the interstitial fluid pressure, and hence increase the penetration of the chemotherapeutic drug into the tumor.\textsuperscript{38,39} The underlying morphological mechanisms have not been clearly elucidated yet. In this study we demonstrate for the first time that decrease in VEGF-A concentration leads to intussusceptive remodeling of the vasculature, including pruning. Combined with our previous study on tumor recovery after antiangiogenic treatment this observation may allude to the mechanism underlying the “normalization” of tumor vasculature: VEGF inhibition not only stops further angiogenesis but also leads to transient tumor shrinkage via vascular elimination associated with vessel normalization by activation of IVP and remodeling.

In conclusion, the mechanism of VEGF-dependent IVP described here gives new insights to the understanding of the process of vasculature regression, which is in line with the newer findings of nonsprouting angiogenesis and might provide new potential targets for antiangiogenic therapy.

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Disclosures
None.

References


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**Supplemental Figure 1.** Verification of the apoptosis of ECs by the investigation of doubled stained (TUNEL and lectin) whole-mount samples harvested at 72 hours after the VEGF application in LSM Zeiss 510 M. The overview picture (a) displays the cross-section through the whole-mount sample with apparent vascular lumen (Lu). The projection image (“Volume”) of a Z-stack of LSM images of endothelial vascular surface (red=endothelial cell surface marker) distinctly shows TUNEL-positive (green) endothelial cells. Nuclei of apoptotic endothelial cells are indicated with asterisks.

**Material and method:**

**Apoptosis (TUNEL) assay combined with endothelial cell surface staining**

The whole-mount samples of the CAM vasculature were harvested and fixed in 2%PFA at 72 hours after the VEGF gel application. They were further stained for fluorescence staining using the TUNEL Label Mix (Roche Diagnostics AG, Switzerland; Cat Nr:11767291910) to visualize the apoptotic cells and rhodamine-labeled Lens culinaris agglutinin (LCA) (Vector Labs, CA, USA; Cat. Nr: RL-1042) for visualization of endothelial cell surface according to the manufacturer’s recommendations. The mounted samples were then investigated in the LSM Zeiss 510M, the obtained Z-stacks of images were afterwards imported in and analysed by Imaris Software.