VE-cadherin is not required for the formation of nascent blood vessels but acts to prevent their disassembly

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We investigated the role of vascular endothelial (VE)–cadherin in blood vessel morphogenesis and established a temporal correlation linking the failure in vessel morphogenesis in VE-cadherin null embryos to a specific step in vasculogenesis. We showed that the sequence in which blood vessels failed followed the order in which they had formed (ie, those forming first—yolk sac, allantoic and endocardial vessels—were the first to display morphologic abnormalities). We next showed that in place of normal reticulated networks of blood vessels, clusters of platelet endothelial cell adhesion molecule–positive (PECAM⁺) cells formed within cultured allantois explants from VE-cadherin null embryos. Similarly, a function-blocking VE-cadherin antibody, BV13, caused PECAM⁺ cell clusters to form in cultured allantois explants from normal mice. Finally, we demonstrated that formation of PECAM⁺ cell clusters in response to BV13 was not due to a disruption in the formation of nascent vessels but was due to the actual disassembly of nascent vessels. Based on these findings, we conclude that the events of de novo blood vessel formation up to the point at which a vascular epithelium forms (ie, nascent vessels with lumens) are not dependent on VE-cadherin and that VE-cadherin, whose expression is up-regulated following vascular epithelialization, is required to prevent the disassembly of nascent blood vessels.

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Introduction

De novo blood vessel formation (vasculogenesis) is the process by which endothelial cell progenitors (angioblasts) interact to form vascular epithelia (ie, nascent blood vessels). As with all epithelia, cadherins play an essential role in mediating vascular epithelial adherens junction formation. Despite intense interest in the functional roles of cadherin family members, the specific roles of cadherins in the formation and regulation of vascular epithelium remains to be elucidated. Endothelial cells express both neural (N)–cadherin¹² and vascular endothelial (VE)–cadherin³ and targeted deletion of these genes in mice leads to early embryonic death with associated severe vascular anomalies.⁴,⁶

A mechanistic understanding of the observed vascular defects in N- and VE-cadherin knockout has not yet been achieved. In N-cadherin–deficient embryos, a number of vascular anomalies were noted, but the anomalies were not attributed to defective vasculogenesis or angiogenesis. In VE-cadherin knockouts, Carmeliet et al⁶ concluded that because lumenized vessels were evident, the basis of the defects was a failure in the process of angiogenesis. By contrast, Gory-Faure et al,⁵ noting defects in the yolk sac vessels concomitant with apparent normal intraembryonic vessels (ie, dorsal aortae), concluded that extraembryonic vasculogenesis was dependent on VE-cadherin activity, whereas intraembryonic vasculogenesis was not.

Herein, we investigated the vascular phenotype of VE-cadherin–deficient embryos to reconcile the disparate views as to whether the protein is required in the process of vasculogenesis. In the first set of studies, we characterized the temporal appearance of vascular anomalies in blood vessel formation occurring in different embryonic tissues of null embryos. In the second series of studies, we used two VE-cadherin monoclonal antibodies, BV13 and BV14, in conjunction with wild-type allantois explant culture to investigate the role of VE-cadherin in the de novo formation of blood vessels. Both BV13, which is directed against the distal extracellular domain of VE-cadherin (EC1), and BV14, which is directed against the more proximal extracellular domain (EC4),⁷ have been shown to have function-blocking activity as judged by their ability to disrupt the formation of cordlike vascular structures in cultured mouse lung endothelial cells⁸ and to disrupt tumor neovascularization in vivo.⁷,⁹

Our findings establish that VE-cadherin function is not critical to the process by which angioblasts transition to form a nascent vascular epithelium. However, because nascent vessels disassemble in the absence of VE-cadherin activity, we reason that VE-cadherin is critically required to maintain or prevent (or both) the disassembly of nascent embryonic blood vessels.

Materials and methods

Antibodies

Purified or fluorescently conjugated antibodies to platelet endothelial cell adhesion molecule (PECAM/CD31, clone Mec-13.3) and Flk1,
fluorescently conjugated isotype control antibodies, and antibodies to VE-cadherin (CD144, clone 11D4.1) were purchased from BD PharMingen (San Diego, CA). Monoclonal antibodies to VE-cadherin, BV13 and BV14, were generated as previously described.\textsuperscript{7,10} Fluorescently conjugated secondary antibodies were purchased from Jackson ImmunoResearch Labs (West Grove, PA).

**Allantois culture**

Allantois were dissected from mouse embryos and cultured as previously described.\textsuperscript{11,12} Four allantois culture schemes were used. (1) To investigate the role of VE-cadherin in vasculogenesis, prevascularized mesoderm of 7.8-days postcoitum (dpc) murine allantois was cultured for a total of 18 hours (37°C; 5% CO\textsubscript{2}) in the presence or absence of experimental reagent. (2) To establish whether VE-cadherin was acting early or late in vasculogenesis, we modified the aforementioned culture scheme and divided the 18-hour culture period into two 9-hour culture periods. Allantois explants were either cultured for 9 hours in the presence of reagent and then cultured an additional 9 hours in serum-containing medium or cultured for 9 hours in serum-containing medium and then exposed to experimental reagent during the last 9 hours of culture. (3) To investigate vascular remodeling associated with vasculogenesis, 8.5-dpc murine allantois were cultured for a total of 18 hours (37°C; 5% CO\textsubscript{2}) in the presence or absence of experimental reagent. (4) Finally, to examine the response of established vessels, 8.5-dpc murine allantois were cultured for 18 hours (37°C; 5% CO\textsubscript{2}) to allow formation of stable vascular networks and then cultured for an additional 24 hours (37°C; 5% CO\textsubscript{2}) in the presence or absence of experimental reagent.

**Fixation, immunolabeling, and microscopy of mouse embryos and cultured allantois**

Procedures for whole-mount immunolabeling of mouse embryos and for immunolabeled cultured allantois have been described previously.\textsuperscript{11,12} Specimens were mounted in an antiphotobleaching mounting medium and all images were acquired at approximately 25°C. Conventional fluorescence or differential interference contrast images of allantois cultures were obtained using a Leica DMR research-grade microscope equipped with Leica objectives (5\times0.15 HC PL Fluotar, 10\times0.30 HC PL Fluotar, 20\times/0.70 HC Plan Apo, 40\times0.85 HC Plan Apo; Leica, Heidelberg, Germany) and a SPOT-RT camera (Diagnostic Instruments, Sterling Heights, MI). Images were acquired using SPOT-RT 3.5.7 software (Diagnostic Instruments). Laser confocal microscopic images were acquired using Zeiss 5\times/0.15 and 10\times/0.30 objectives (Zeiss, Thornwood, NY) on a Bio-Rad MRC 1024 laser-scanning confocal microscope (Bio-Rad, Microscopy Division, Cambridge, MA) equipped with Lasersharp 2000 software (Bio-Rad Cell Science Division, Hemel Hempstead, United Kingdom). Confocal z-series were projected using Image J 1.31v (National Institutes of Health, Bethesda, MD). Montages of microscopic images were generated using Adobe Photoshop 7.0 software (Adobe Systems, San Jose, CA).

**Fluorescence-activated cell scanning**

Cultured allantois were dissociated in 1× trypsin-EDTA (ethylenediaminetetraacetic acid) solution (40 allantois/0.5 mL) for 7 minutes at 37°C. Dulbecco modified Eagle medium (DMEM) containing 10% fetal bovine serum (FBS) and 2% penicillin/streptomycin/L-glutamine (PS-G; 0.5 mL) was added to inhibit the trypsin. The cell suspension was centrifuged at 1100g for 10 minutes using an IEC Centra CL2 benchtop centrifuge (IEC, North Hempstead, MA). The cell pellet was resuspended in 0.5 mL DMEM containing 10% FBS and 2% PS-G and incubated for 40 minutes at 37°C. The cell suspension was resuspended in 1% bovine serum albumin (BSA), 0.01% NaN\textsubscript{3}, and 10 ng/mL DNase (fluorescence-activated cell scanning [FACS] buffer) and pelleted by centrifugation as before. Cells were counted on a hemocytometer, separated into tubes for immunolabeling (final concentration of 1×10\textsuperscript{6} cells/mL), and immunolabeled with fluorescently conjugated isotype control antibodies, fluorescein isothiocyanate (FITC)–conjugated PECAM, phycoerythrin (PE)–conjugated Flk-1, or both for 40 minutes on ice. Antibodies were added to the cell suspensions at a final concentration of 100 ng/1×10\textsuperscript{6} cells and incubated 40 minutes at 4°C. Cells were pelleted and washed in FACS buffer. The washed cells were resuspended in FACS buffer containing 0.1 mg/mL propidium iodide (Sigma, St Louis, MO). After propidium iodide treatment the cells were washed in FACS buffer, centrifuged, resuspended in 0.2 mL FACS buffer, and subjected to FACS analysis using a FACScalibur (Becton Dickinson, San Jose, CA).

**Results**

**Temporal failure of vasculogenesis in VE-cadherin null embryos**

The presence of both normal and abnormal blood vessels in the VE-cadherin nulls, as described by Gory-Faure et al.,\textsuperscript{5} led us to hypothesize that the vascular defects might correlate with the temporal order in which blood vessels are formed de novo in the mouse embryo. This hypothesis is based on our previous findings\textsuperscript{11} showing that vasculogenesis is initiated regionally at different times during development with extraembryonic vessels being the first to form (yolk sac and allantoic vessels) followed by the endocardium, dorsal aortae/sinus venosus, and the cranial vessels. Our analysis of VE-cadherin null embryos showed that vessels of the allantois and endocardium were abnormal in 8.5-dpc embryos, whereas the dorsal aortae in these embryos...
appeared normal (Figure 1). Further support for a temporal failure in blood vessels in the VE-cadherin nulls was the finding that the dorsal aortae, which appeared normal at 8.5 dpc, were abnormal in 9.5-dpc null embryos (Figure 1F,I). Based on these observations and the temporal order for the formation of specific embryonic vascular networks (yolk sac, allantois, endocardium, dorsal aortae/sinus venosus, and cranial vasculature), it was evident that the vascular networks that had developed first (ie, yolk sac and allantoic vessels) were the first to fail. Vessels that form later in development, such as the dorsal aortae, failed at a later stage in development.

**VE-cadherin–null allantois explants fail to undergo normal vascular morphogenesis in culture**

To further assess the role of VE-cadherin in the temporal events of vascular morphogenesis, we turned to the allantois explant culture model. At 8.5 dpc, the allantois has an extensive network of blood vessels (Figure 2A). When 8.5-dpc allantois explants are cultured for 18 hours, the explants flatten and the network of PECAM⁺ capillary-like vessels is evident within the adherent explants (Figure 2B). Following 18 hours of culture, allantois explants from 8.5-dpc VE-cadherin nulls failed to develop interconnected networks of PECAM⁺ cells as compared to wild-type controls (compare Figure 2C and 2D). Instead, only isolated clusters of PECAM⁺ cells were present (Figure 2D).

**VE-cadherin function-blocking antibody treatment phenocopies vascular anomalies observed in genetic VE-cadherin deficiency**

We next evaluated the role of VE-cadherin in vascular morphogenesis using function-blocking VE-cadherin monoclonal antibodies in conjunction with the allantois explant culture model. Treatment of allantois explants from normal 8.5-dpc mouse embryos with BV13 disrupted vascular morphogenesis (Figure 3B). In place of interconnected vessels that are present in control cultures (Figure 3A), only PECAM⁺ clusters of cells were evident in the treated cultures (Figure 3B). By contrast, BV14 treatment produced little or no effect on vascular morphogenesis occurring in cultured 8.5-dpc allantois (Figure 3C). The fact that BV13 treatment produced PECAM⁺ aggregates in 8.5-dpc allantois explants initially possessing extensive vascular networks (Figure 2A) suggested that the antibody was acting to promote blood vessel disassembly.

**Established vessels formed in long-term cultures of normal 8.5-dpc allantois exhibit incremental disassembly in response to BV13 treatment**

To substantiate the hypothesis that BV13 antibody was promoting the disassembly of blood vessels, we evaluated its effects on established vessels within 8.5-dpc explants cultured for 18 hours prior to administration of the antibody. Within 6 hours after BV13 treatment, allantois vascular networks had discontinuities in the normally interconnected vascular network (Figure 4C). Following 12 hours of BV13 treatment, networks were absent and only clusters of PECAM⁺ cells were apparent (Figure 4D). Taken together, these results support the hypothesis that VE-cadherin is necessary for the stability of vascular networks that are present at 8.5 dpc and suggest that VE-cadherin functions throughout vascular development to stabilize blood vessel networks.
together, these findings suggest that VE-cadherin function is required to prevent disassembly of nascent embryonic blood vessels and that in the absence of VE-cadherin activity nascent blood vessels incrementally disassemble over a 12-hour period.

VE-cadherin antibody-mediated blood vessel disassembly does not involve alteration of mesodermal or endothelial subpopulations

We next used flow cytometry to evaluate the effects of BV13 antibody on subpopulations of cells within the allantois. The analysis revealed that BV13 antibody treatment did not elicit significant changes in either the number of angioblasts (Flk1⁺/PECAM⁻ population) or endothelial cells (Flk1⁺/PECAM⁺ population) present in 8.5-dpc explant cultures (Figure 5). Furthermore, the antibody treatment did not influence the number of dead cells in the explant cultures as measured by propidium iodide exclusion. These findings suggest that blood vessel disassembly induced by BV13 treatment is not the result of significant levels of cell death.

VE-cadherin function-blocking antibody treatment disrupts vascular morphogenesis in the 7.8-dpc allantois explant culture model

In addition to establishing the role of VE-cadherin in preventing the disassembly of newly formed blood vessels, we also evaluated its potential role in the process of de novo blood vessel formation. To do this we used the 7.8-dpc allantois explant culture model in which prevascularized allantois explants are obtained from 7.8-dpc embryos and cultured for 18 hours. During this culture period, blood vessels form de novo (Figure 6A). As shown in Figure 6B, introduction of BV13 at the initiation of the culture period had profound consequences on vascular morphogenesis. In contrast to control explants (Figure 6A) in which interconnected PECAM⁺ networks of vessels were evident, only clusters of PECAM⁺ cells could be observed in the BV13-treated cultures (Figure 6B). High-magnification fluorescence imaging of the BV13-treated explants shows that the clusters are composed of small numbers of cells with PECAM immunolabeling in cell-cell junctions (compare arrows in control, Figure 6C, to BV13 treated, Figure 6D). The effects of BV13 on vascular morphology can also be observed using differential interference contrast (DIC) microscopy (compare arrows in control, Figure 6E, to BV13-treated, Figure 6F). In separate experiments, we showed that Fab fragments of BV13 antibody (24 μg IgG/mL; G) or BV14 antibody (24 μg IgG/mL; H) had little or no effect on vasculogenesis in cultured 7.8-dpc explants (Figure 6H).

Figure 5. VE-cadherin antibody-mediated blood vessel disassembly does not involve alteration of mesodermal or endothelial subpopulations. (A-B) Representative FACs profiles of cells isolated from control and BV13-treated (24 μg IgG/mL) 8.5-dpc cultured allantois, respectively. (C) A graphic compilation of the FACs data from 4 experiments comparing control versus BV13-treated cultures. Together these analyses show no significant changes in the Flk1⁺ (angioblasts; filled bars), Flk1⁺/PECAM⁺ (endothelial cells; dark gray bars), or propidium iodide⁺ (dead cells; light gray bars) cell populations in BV13-treated allantois cultures as compared to control cultures. Error bars indicate the standard deviation of the mean.

Figure 6. VE-cadherin function-blocking antibody treatment disrupts vascular morphogenesis in the 7.8-dpc allantois explant culture model. (A-D) PECAM-immunolabeled allantois explants from 7.8-dpc embryos following 18 hours of culture in control medium (A,C) or BV13 monoclonal antibody (24 μg IgG/mL; B,D). Arrows (C, control; D, BV13-treated) indicate PECAM deposition at cell-cell junctions. (E-F) DIC images of the same regions of the explants depicted in panels C and D, respectively. Arrowheads in panels D,F indicate a PECAM⁺ cluster of cells resulting from BV13 treatment. (G-H) PECAM-immunolabeled allantois explants from 7.8-dpc embryos following 18 hours of culture between fragments of BV13 antibody (24 μg IgG/mL; G) or BV14 antibody (24 μg IgG/mL; H). Bars equal 50 μm. Panels A, B, G, and H are confocal images; C and D, epifluorescence images; and E and F, DIC images.

Initial events of vasculogenesis including the formation of nascent endothelial tubes are not dependent on VE-cadherin activity

If the phenotype observed in BV13-treated allantois (ie, clusters of PECAM⁺ cells) was due to blood vessel disassembly resulting from the loss/lack of VE-cadherin function, we reasoned that the antibody should, therefore, not perturb earlier events in vasculogenesis (ie, the formation of nascent blood vessels). To test this hypothesis, 7.8-dpc allantois explants were cultured in the presence of BV13 antibody for varying periods of time. As shown in Figure 7C, blood vessel formation in 7.8-dpc
allantois explants was not perturbed by treatment with BV13 for 9 hours from initiation of the culture. By contrast, when 7.8-dpc allantois explants were cultured for 9 hours in the absence of antibody and then exposed to BV13 for an additional 9 hours, PECAM⁺ clusters of cells replaced the normal reticulated vascular networks (Figure 7D).

Discussion

After globally assessing embryonic blood vessel formation in VE-cadherin null embryos, we established that the regional vascular abnormalities could be correlated with the age of the given vascular network. The temporal order of the formation of discrete vascular networks/blood vessels in normal mouse embryos is yolk sac and allantoic vessels, the endocardium, dorsal aortae/sinus venosus, and the cranial vascular network. Vessels that were first to form in VE-cadherin null embryos (ie, yolk sac and allantois vessels) were the first to display morphologic abnormalities. Our findings in this regard are consistent with observations by Gory-Faure et al, who also noted aberrant yolk sac vessels in VE-cadherin null embryos (ie, yolk sac and allantois vascular morphogenesis when VE-cadherin function is essential). For example, administration of either neutralizing antibodies to embryonic vessels formed as part of vasculogenesis (ie, they are nascent endothelial tubes), the disassembly effect that we observed may be the underlying basis for the capacity of BV13 and other VE-cadherin function-blocking antibodies to inhibit tumor vascularization in vivo. In this context, our findings that BV13 but not BV14 promotes the disassembly of vessels serves to support the conclusions of Corada et al, that the mechanism by which these two antibodies inhibit tumor neovascularization differ.

The clusters of PECAM⁺ cells that form in response to either genetic VE-cadherin deficiency or functional inhibition by VE-cadherin antibody are similar to the clusters of PECAM⁺ cells that result from coalescence of angioblasts during normal vasculogenesis and the clusters of PECAM⁺ cells that result from inhibition of vascular endothelial growth factor (VEGF) signaling during vasculogenesis. For example, administration of either neutralizing antibodies against recombinant mouse VEGF164 or the soluble form of VEGF receptor-1 (sFlt1) to avian embryos or cultured murine allantois explants results in PECAM⁺ cell clusters. The similarity between the response to VE-cadherin deficiency and suppression of VEGF signaling suggests a link that is critical to vasculogenesis. Numerous studies support this possibility, including reports that VEGF receptor-2 (Flk1) and VE-cadherin can be coprecipitated from endothelial cell preparations. VE-cadherin deficiency reduces the half-life of Flk1, and VEGF can induce VE-cadherin tyrosine phosphorylation in endothelial cells, and VE-cadherin–deficient endothelial cells are unresponsive to VEGF signaling. However, the coalescence of angioblasts that normally occurs during early vasculogenesis, as well as the transition of angioblast aggregates to a vascular epithelium, involves cells that express Flk1 but not VE-cadherin. In the yolk sac, allantois and
embryo proper, VE-cadherin expression is up-regulated between 8 and 8.5 dpc, 20 stages when PECAM+ endothelia are detectable in each of these regions. Therefore, we can exclude a role for an integrated Flk1–VE-cadherin signaling pathway up to the point in which lumenized blood vessels are formed, the pre-epithelial phase of vasculogenesis. With the onset of VE-cadherin expression, integration of Flk1 signaling and VE-cadherin function is possible. The functional significance of this integration is likely to involve regulation of cell-cell interaction and permeability in response to VEGF.

The VE-cadherin knockout mouse teaches us that blood vessels can form in the absence of VE-cadherin. 5, 6 Herein we show that perturbation of VE-cadherin leads to disassembly of nascent vessels. These findings seem incongruent unless one evokes a role for VE-cadherin in which it serves to regulate the disassembly of vessels. In other words, its expression in established blood vessels imparts a mechanism by which cell-cell adhesion can be diminished to allow permeability. Such a role is also consistent with the mechanism of action of VEGF to regulate vascular permeability. Therefore, the integration of Flk1 signaling and VE-cadherin function confers on the endothelium the ability to modulate cell-cell adhesion in response to VEGF.

Assigning a role for VE-cadherin in either vasculogenesis or angiogenesis is dependent on one’s definition of these processes. Although there is a relatively clear understanding of many of the steps in the process of vasculogenesis (ie, the birth of endothelial progenitors [angioblasts], coalescence of angioblasts into cordlike structures, and lumenization to form a vascular endothelium), a clear end point of the process is lacking. One end point may be when mural cells have invested the blood vessel. In the allantois, smooth muscle cell antigens (ie, smooth muscle α actin) are not detectable until 9.0 dpc (P.A.F., unpublished observations, September 2003). The fact that vascular anomalies are apparent in the allantoides of 8.5-dpc VE-cadherin nulls and the fact that VE-cadherin function-blocking antibodies have dysmorphogenic effects on the vasculature of 7.8-dpc allantoic explants indicates that VE-cadherin expression and therefore function is essential at the point following the formation of vascular endothelium but prior to mural cell investment. Based on this point of view, VE-cadherin is critical for vasculogenesis.

Acknowledgment

The authors thank Haiqun Zeng for her expert technical assistance with fluorescence-activated cell scanning.

References

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