An in vivo model allowing continuous observation of human vascular formation in the same animal over time

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Angiogenesis contributes to numerous pathological conditions. Understanding the molecular mechanisms of angiogenesis will offer new therapeutic opportunities. Several experimental in vivo models that better represent the pathological conditions have been generated for this purpose in mice, but it is difficult to translate results from mouse to human blood vessels. To understand human vascular biology and translate findings into human research, we need human blood vessel models to replicate human vascular physiology. Here, we show that human tumor tissue transplantation into a cranial window enables engraftment of human blood vessels in mice. An in vivo imaging technique using two-photon microscopy allows continuous observation of human blood vessels until at least 49 days after tumor transplantation. These human blood vessels make connections with mouse blood vessels as shown by the finding that lectin injected into the mouse tail vein reaches the human blood vessels. Finally, this model revealed that formation and/or maintenance of human blood vessels depends on VEGFR2 signaling. This approach represents a useful tool to study molecular mechanisms of human blood vessel formation and to test effects of drugs that target human blood vessels in vivo to show proof of concept in a preclinical model.

Angiogenesis, the development of new blood vessels from preexisting vasculature, is essential under many physiological and pathological conditions. Dysregulated angiogenesis contributes to numerous malignant, ischemic, inflammatory, infectious and immune diseases. While an understanding of blood vessel formation is needed to treat these diseases, vascular development and maintenance is a complex process that is difficult to recapitulate in vitro. Because of this, in vivo models are needed for the study of vascular change and drug development. Experimental mouse models that better represent pathological conditions have been generated for a improved understanding of angiogenesis and therapy. However, it is difficult to translate experimental results from mouse to human blood vessels. For example, a single cell analysis revealed that endothelial cells are genotypically heterogeneous, and human and mouse endothelial cells have different molecular expression patterns. A previous report showed that tumor xenografts vascularized with human blood vessels grow more rapidly and are more resistant to chemotherapeutic or anti-angiogenic drugs than are murine blood vessels. Thus, human blood vessel models are required to understand human vascular biology and to bridge preclinical testing to clinical trials for drug development.

Recently, several in vivo models of human blood vessels have been developed using both human endothelial cells (ECs) and human tissues. These models include transplanting human umbilical vein endothelial cell (HUVECs) spheroids subcutaneously, mesenchymal precursor cells and EC-derived blood vessels or human stem cell-derived vascular organoids. Although human blood vessels were detected in these human tumor tissue transplant models, they are rapidly replaced by murine blood vessels. Furthermore, in such transplantation models, detection of human tumor blood vessels has been accomplished on the basis of immunohistochemistry and/or immunofluorescence using sections of harvested tissues. These methods provide a snapshot of a dynamically changing system of spatial interactions and molecular landscapes; however, these methods cannot provide images of the kinetics of changes to the vasculature within the tissue of the same individual mouse. To solve this problem, in vivo imaging is a useful method because this technique allows continuous observation of the blood vessels in a non-invasive manner. Recently, Meehan et al. developed a human subcutaneous fat

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transplantation model to visualize vascular cytoadherence of erythrocytes infected with the human parasite Plasmodium falciparum over a short period of time. While tissue-derived human blood vessel engraftment has been successful, to the best of our knowledge, continuous observation of human tissue-derived blood vessels for an extended period of time has not yet been reported.

In the current study, to develop a model allowing the continuous observation of human tissue-derived blood vessels, we used a human tumor tissue transplantation approach because tumor blood vessels have highly angiogenic potential. We chose the cranial window model to monitor human blood vessels because this uses skull bone for fixing the cover glass to enable a stable rigid fixation allowing long-term observation for up to a year. Using this method, we were able to observe human blood vessels in tumor tissues over the long term, for at least 49 days; thus, given the lack of cancer cell growth in our model, we were able to investigate human angiogenesis in the absence of tumor cells. Here, we have assessed whether anti-human VEGFR2 antibody affects blood vessel formation in this model and we discuss its suitability for use in drug screening for new modulators of human angiogenesis by allowing observations on three dimensional blood vessel structures over time.

Methods

Ethics statement. Informed consent was obtained from patients for the use of their tumor tissues for transplantation. This study was conducted under recognized ethical guidelines (Declaration of Helsinki) and approved by Osaka University Institutional Review board. Human tumor tissues were distributed by a human tissue bank (the JCRB Bioresource Bank, National Institutes of Biomedical Innovation, Health and Nutrition, Osaka, Japan). The clinical characteristics of the four human tumor tissues are provided in Table 1.

Mice. NOD.CB17-Prkdc<sup>scid</sup>/Jcl (NOD-Scid) mice were purchased from CLEA Japan (Tokyo, Japan). All animals were housed in a temperature-controlled environment with a 12 h light/dark cycle, and food and water was freely provided. All mouse experiments were performed under the guidelines of the Osaka University Committee for Animal and Recombinant DNA Experiments (Approval number 4062).

Transplantation of human tumor tissues into the cranial window. An overview of the methodology used to generate the xenograft model is shown in Fig. 1. Freshly isolated human tumor tissue was washed...
twice with HBSS (Thermo Fisher Scientific, Waltham, MA) containing 100 μg/ml kanamycin sulfate (FUJI-FILM Wako Pure Chemical Corporation, Osaka, Japan), and 0.5 μg/ml amphotericin B (Thermo Fisher Scientific). It was then was shipped within 2 h after surgery in HBSS containing kanamycin, amphotericin B. On arrival, tumor tissue was washed × 4 with saline (Otsuka Pharmaceutical Factory, Tokushima, Japan) and then cut into 1–2 mm pieces in DMEM (high glucose, no glutamine, no phenol red) (Thermo Fisher Scientific) containing 10% heat-inactivated-FBS (Sigma-Aldrich), 100 units/mL penicillin/100 μg/mL streptomycin (p/s, Sigma-Aldrich). Tumor fragments were placed in HBSS containing 100 ng/mL recombinant human VEGF165 (ProTec, Rocky Hill, NJ), 100 ng/mL recombinant human bFGF (Peprotech), and p/s. NOD-Scid mice were anesthetized with three mixed anesthetic agents including 0.3 mg/kg of medetomidine (Domitor, Nippon Zenyaku Kogyo, Fukushima, Japan), 4.0 mg/kg of midazolam (Sandoz, Holzkirchen, Germany), and 5.0 mg/kg of butorphanol (Vetorpharm, Meiji Seika Pharma, Tokyo, Japan). To prevent cerebral edema during surgery, mice were injected intraperitoneally with 0.2 mg/kg dexamethasone sodium phosphate (Dexart, Fuji Pharma Co., Tokyo, Japan), 5 mg/kg carprofen (Rimadyl, Zoetis, Parsippany-Troy Hills, NJ), 300 μL 10% glycine and 5% fructose (Glycele, Chugai Pharmaceutical, Tokyo, Japan). Holes were drilled in the skulls of anesthetized animals to expose the dura mater by slowly lifting the central islands of skull bone. The human tumor fragments were transplanted onto the dura mater. The transplanted areas were filled with artificial cerebrospinal fluid (Artcereb, Otsuka Pharmaceutical Factory) and sealed with a silicon-coated 5 mm round glass cover slip (Matsunami Glass Ind., Osaka, Japan) using a medical adhesive (Aron Alpha A, Daiichi Sankyo, Tokyo, Japan). Mice were kept on a heated pad set to 37 °C until adhesion between glass and skull bone was complete, and then the anesthetic was reversed with 0.3 mg/kg of atipamezole (Antisedan, Nippon Zenyaku Kogyo). To transplant tumor tissues at different time points, samples were preserved in Cellbanker 1plus (Takara Bio, Shiga, Japan) until use.

In vivo visualization of human blood vessels. In vivo visualization of human blood vessels was performed with a TCS/SP8 (Leica Microsystems, Wetzlar, Germany) equipped with a Chameleon laser (Coherent, Santa Clara, CA). To specifically visualize the human blood vessels, Alexa Fluor 488-conjugated anti-human CD31 monoclonal antibody (huCD31-4F88 Ab, clone: MW59, 303110, Biolegend, San Diego, CA) was injected intravenously. The huCD31-4F88 antibody was allowed to circulate for 7 days and establish stable fluorescence before imaging started. To study in vivo blood flow through the human blood vessels, mice were injected with 1 mg 70 kD Rhodamine B isothiocyanate-dextran (Sigma-Aldrich) dissolved in 100 μL PBS and then observed by microscopy. During the imaging, the animal's head was immobilized using a custom-made cranial window holder. Mice were kept on a heated pad set to 37 °C, and anesthesia was maintained with isoflurane vaporizer and scavenger (Muromachi Kikai, Tokyo, Japan). To acquire high-resolution images of whole transplanted human blood vessels, these regions were subdivided into multiple (nine to eleven) smaller images. These subdivided regions were imaged individually and then combined via stitching to a larger overview.

Transrentizing tissues with the dura mater. The transparentizing solution BABB is a solution of 1 part benzoic acid (FUJIFILM Wako Pure Chemical) plus 2 parts benzyl benzoate (FUJIFILM Wako Pure Chemical). Whole-mount tissue samples were stained with Armenian hamster anti-mouse CD31 mAb (MAB198Z, Merck Millipore, Darmstadt, Germany, dilution 1/1000), and then stained with Alexa Fluor 647-conjugated anti-Armenian hamster pAb (127-605-160, Jackson ImmunoResearch Laboratories, West Grove, PA, dilution 1/1000). Tissues were treated with 50% methanol/PBS, 100% methanol, 50% BAB/Methanol twice for 10 min each and with 100% BABB over night.

Ramucirumab treatment. Mice with successfully engrafted human vessels were divided into two groups, one for ramucirumab treatment and one for the isotype IgG control. Fourteen days after tumor transplantation, mice were treated with 40 mg/kg ramucirumab (Cynamza, Eli Lilly and Co., Indianapolis, IN) or isotype IgG from human serum (I4506, Sigma-Aldrich) intraperitoneally three times per week.

Immunofluorescence staining. Tissue fixation and staining of sections or tissues with antibodies was performed using the following method. Excised dura mater was fixed with 4% paraformaldehyde (PFA) in PBS for four hours. It was then embedded in FSC22 Blue (Leica Microsystems) and frozen in liquid nitrogen. Subsequently, 14 μm cryosections were obtained via the Kawamoto film method16. An Armenian hamster anti-CD31 monoclonal antibody (C6198, Sigma-Aldrich) was injected subcutaneously for four hours. It was then embedded in FSC22 Blue (Leica Microsystems) and frozen in liquid nitrogen. Subsequently, 14 μm cryosections were obtained via the Kawamoto film method16. An Armenian hamster anti-α-smooth muscle actin (αSMA) mAb (C6198, Sigma-Aldrich, dilution 1/100), rabbit anti-human cytokeratin 19 mAb (ab52625, Abcam, Cambridge, UK, dilution 1/100), Alexa Fluor 546-conjugated goat anti-rabbit IgG pAb (A-11010, Thermo Fisher Scientific, dilution 1/200), Alexa Fluor 546-conjugated streptavidin (S32357, Thermo Fisher Scientific, dilution 1/400), Cy3-conjugated mouse anti-human nuclei (HuNu) mAb (MAB1281B, Merck Millipore, 1/400), Cy3 conjugated mouse anti-a-smooth muscle actin (aSMA) mAb (C6198, Sigma-Aldrich, dilution 1/100), rabbit anti-cytokeratin 19 mAb (ab52625, Abcam, Cambridge, UK, dilution 1/100), Alexa Fluor 488-conjugated anti-rat IgG pAb (A-11018, Thermo Fisher Scientific, dilution 1/200), Alexa Fluor 488-conjugated goat anti-rabbit IgG pAb (A-11010, Thermo Fisher Scientific, dilution 1/200), Alexa Fluor 647-conjugated streptavidin (S32357, Thermo Fisher Scientific, dilution 1/400), Cy3-conjugated goat anti-Armenian hamster IgG pAb (127-605-160, Jackson ImmunoResearch Laboratories, dilution 1/100), and Alexa Fluor 647-conjugated goat anti-Armenian hamster IgG pAb (127-605-160, Jackson ImmunoResearch Laboratories, dilution 1/400) were used for stain- ing. Cell nuclei were visualized with TO-PRO-3 (T3605, Thermo Fisher Scientific, dilution 1/1000) or Hoechst 33342 (Sigma-Aldrich, 1 μg/mL). To avoid non-specific streptavidin/biotin stain, tissues were blocked using the streptavidin/biotin blocking kit (Vector Laboratories, Burlingame, CA) according to the manufacturer’s protocol.

Image acquisition and analysis. Immunofluorescence images were obtained with confocal microscopy (TCS/SP8 or TCS/SP5; Leica Microsystems). At least 3 images per sample were acquired. Images were processed
using the Leica application suite (Leica Microsystems) and Adobe Photoshop CC software (Adobe, San Jose, CA). To quantify the length of blood vessels, CD31-positive immunofluorescence images were analyzed with the computational analysis tool Angiotool17. Each calculated value was plotted graphically.

Cell culture. MS-1 (mouse pancreatic islet endothelial cell line) was purchased from the ATCC (Manassas, VA) and maintained in DMEM (Sigma-Aldrich, St. Louis, MO) supplemented with 5% heat-inactivated-FBS and p/s. HUVECs (human umbilical vein endothelial cells) were purchased from Kurabo (Osaka, Japan) and maintained in HuMedia-EB2 supplemented with HuMedia-EG (Kurabo).

Flow cytometric analysis. HUVECs and MS-1 cells were harvested by trypsinization and incubated in 4% FCS PBS containing human Fc receptor blocking solution (Biolegend, dilution 1/1000) or mouse Fc receptor blocking solution (Becton, Dickinson and Co., Franklin Lakes, NJ, dilution 1/1000) for 5 min at room temperature. These cells were incubated with anti-huCD31-AF488 Ab (dilution 1/200) and Alexa Fluor 647-conjugated mouse CD31 antibody (clone: MEC13.3, 102516, Biolegend, dilution 1/200) in 4% FCS PBS at 4 °C for 30 min. After washing with 4% FCS PBS twice, cells were resuspended in 300 µL 4% FCS PBS containing SYTOX Blue (Thermo Fisher Scientific, dilution 1/1000) and analyzed on a BD LSRFortessa (Becton Dickinson). Data were analyzed using FACS Diva software and FlowJo version 10.6.2 (Becton Dickinson).

Statistical analysis. Data are shown as mean ± SD. Statistical analysis was conducted by a Welch t test or one-way ANOVA with Dunnett’s multiple comparison test using GraphPad Prism 7 statistical software (GraphPad Software, San Diego, CA). A p-value of < 0.05 was judged as significant.

Results

Confirmation of specific recognition of human ECs. To distinguish human from mouse blood vessels in vivo, we tested the commercially available fluorescently-labeled antibody CD31 that recognizes human ECs14, and confirmed its species specificity using human ECs HUVECs and mouse ECs MS-1 cells. We tested Alexa Fluor 488-conjugated human CD31 antibody (huCD31-AF488 Ab) and Alexa Fluor 647-conjugated mouse CD31 antibody (moCD31-AF647Ab). Flow cytometric analysis showed that huCD31-AF488 Ab detected the human CD31 extracellular domain and moCD31-AF647 Ab detected the mouse CD31 extracellular domain specifically (Fig. 2a,b). In addition, we confirmed the specificity of the antibody by immunocytochemistry (Sup-
Next, to determine whether huCD31-AF488 Ab recognizes human blood vessels, we performed immunofluorescence studies on human colorectal tumor tissues. The huCD31-AF488 Ab recognized tumor blood vessels in colorectal tumor (Fig. 2c). Taken together, these results show that huCD31-AF488 Ab allowed the specific observation of human blood vessels.

In vivo observation of human tumor-derived human blood vessels in the cranial window.

Previous reports showed that transplanted human tumor-derived xenografts initially contained human blood vessels12,18; however, immunohistochemistry showed that intra-tumor human blood vessels decreased markedly 3 days after transplantation or after in vivo passage to secondary recipient mice. In order to observe the change of human blood vessels in our model, we transplanted human tumors into cranial windows and continuously observed human-specific tumor blood vessels by injecting huCD31-AF488 Ab intravenously. Whole-mount staining showed that huCD31-AF488 Ab allowed visualization specifically of human blood vessels in vivo (Fig. 3a).

Figure 3. Human tumor-derived human blood vessels in cranial windows. (a) Representative image of colon tumor patient-derived human blood vessels (green, huCD31-AF488 Ab injected intravenously) and mouse blood vessels (magenta, anti-moCD31 immunofluorescence staining) in the cranial window. White dashed line shows the edge of the engrafted human tissues on day 19. (b) Quantification of total human blood vessel length two weeks after transplantation into cranial the window (Patient #1: n = 8, Patient #2: n = 7, Patient #3: n = 9, Patient #4: n = 4). (c) In vivo perfusion of human blood vessels shown by intravenous injection of Rhodamine B-conjugated 70 kDa Dextran (DextranRhodamine B, magenta) in the xenograft. Image acquisition was at day 20 after tumor transplantation. Lower images show high magnification of the area indicated by dashed boxes in the upper row. White arrowheads show perfused human blood vessels. (d) Perfusion of human blood vessels (green, huCD31-AF488 injected intravenously) in the cranial window was assessed by injecting Rhodamine-conjugated Ulex europaeus agglutinin I (magenta, UEA-Irhodamine). (e) Quantification of perfused human blood vessels in the cranial window (Patient #1: n = 2, Patient #2: n = 2).

We transplanted four human tumors (3 colorectal tumors, and one gastric tumor) into cranial windows. After two weeks, in vivo imaging showed that human blood vessels from different tumor patients were engrafted (Supplementary Fig. S2a–c). We examined the relationship of engrafted human blood vessels and the original tumor blood vessels (Supplementary Fig. S2d). However, we could not identify any factors that influenced the engraftment of human blood vessels in this model. Next, to determine whether transplanted human blood vessels were functional, we injected Rhodamine B-conjugated 70 kDa Dextran into the tail vein. In vivo imaging showed that human blood vessels were partially perfused (Fig. 3c). To further investigate this, we injected the Rhodamine-conjugated human blood vessel-specific lectin, Ulex europaeus agglutinin I (UEA-Irhodamine)19. The degree of perfusion through the human blood vessels was different in grafts derived from the different patients (Fig. 3d,e). These results show that tumor-derived human blood vessels survive in a cranial window model and are partially still perfused after 2 weeks.

Continuous increase of tumor-derived human blood vessels. The cranial window model allows continuous observation of blood vessels over time in the same animal in vivo6. As seen in Fig. 4a,b, the length of
human blood vessels significantly increased from day 21 to day 49, which is inconsistent with results reported in a previous study. In addition, the number of human vessel junctions and average vessel diameter increased (Supplementary Fig. S3a,b). Vessel density, on the other hand, was unchanged (Supplementary Fig. S3c). Immunofluorescence analysis showed that parts of the human blood vessels are positive for Ki-67 (Fig. 4c). Moreover, Ki-67 positivity was significantly higher in the human than mouse blood vessels (Fig. 4d). These results suggest that proliferation of ECs was induced predominantly in the human ECs in our model. Because transplanted human blood vessels have an immature structure, we determined whether they were covered with α-smooth muscle actin (αSMA)-positive pericytes. The results indicated that the human blood vessels were indeed partially covered with αSMA-positive pericytes (percent of pericyte coverage: 13.8% ± 10.2%) (Supplementary Fig. S3d,e). Tumor angiogenesis is facilitated by factors secreted by cancer cells and the surrounding stroma cells, for example vascular endothelial factors (VEGFs) and fibroblast growth factor (FGFs). However, there were no detectable colorectal cancer cells in this model (Supplementary Fig. S3f,g). Thus, these results indicate that tumor-derived human blood vessels have angiogenic potential in the cranial window and that angiogenic signals are derived from non-cancer cells.

The growth of human blood vessels depends on VEGFR2 signaling. Even though several angiogenic factors have been identified, VEGF is one of the most important molecular cues for the formation of new blood vessels. To examine whether growth of tumor-derived human blood vessels in our cranial window model also depends on VEGF signaling pathways, the anti-human VEGFR2 antibody ramucirumab was given three times a week from day 14 to day 21 after tumor transplantation (Fig. 5a). Ramucirumab induced a rapid decrease of human blood vessels (Fig. 5b–d, Supplementary Fig. S4a–c). To determine whether ramucirumab affects only human but not mouse blood vessels, tumor-transplanted dura mater was retrieved on day 21 after transplantation. Ramucirumab treatment decreased human blood vessels but did not inhibit the formation of mouse blood vessels (Fig. 5e–g). It is generally accepted that low-dose anti-angiogenic therapy induces vascular normalization. To test whether ramucirumab does induce vascular normalization, we examined αSMA pericyte coverage. However, we found that ramucirumab did not influence αSMA pericyte coverage in our model (Supplementary Fig. S4d,e). These results show that ramucirumab inhibits VEGF2 signaling of human blood vessels. Taken together, our data show that human blood vessels in cranial windows depend on VEGFR2 signaling.
Discussion

To evaluate human blood vessel formation in vivo, most earlier studies compared tumor sections from different mice by immunohistochemistry or immunofluorescence\textsuperscript{11,12}. However, these methods only show a snapshot of human blood vessel presence at the time that the tissue was obtained. Here, we developed a method for observing human blood vessels continuously over time in the same animal. This model of tumor transplantation into a cranial window directly revealed changes to human blood vessels by using in vivo fluorescence imaging techniques.

Previous studies reported no staining with the anti-human CD34 antibody in the colorectal tumor xenograft model on day 30, and that human tumor blood vessels had been replaced by mouse blood vessels\textsuperscript{11}. In our current model, human blood vessels remained stable and functional for at least 49 days in vivo. We found that some of the ECs in the tumor had high proliferative potential; thus, given the lack of cancer cell growth in our model, human blood vessels are maintained depending on non-cancer cell-derived VEGF in this cranial window model. Further analysis of the origin of this VEGF is required; however, it is apparent that our method differs significantly from previous reports on the growth of human blood vessels in vivo. In a recent study, Koh et al. showed that dura mater ECs had more angiogenic potential than brain ECs, and that VEGFR2 signaling in the dura mater promoted vascular regeneration in an experimental head injury model\textsuperscript{23}. Considering these results, damage to the dura mater caused by tumor transplantation may promote VEGF-dependent EC sprouting. We

\textbf{Figure 5.} VEGFR2 inhibition causes regression of human blood vessels. (a) Schema for tumor transplantation into the cranial window and ramucirumab or isotype IgG treatment. b-d, VEGFR2 inhibition decreases human blood vessels. Representative images of huCD31-AF488-positive human blood vessels (huCD31, green) (b). Quantification of total human blood vessel length (c) and relative length (d) after ramucirumab (Ram) or isotype IgG (Isotype) treatment. e-g: Transplanted tissues were sectioned for the analysis of human and mouse blood vessels. Representative images for the expression of human CD31 (huCD31, green), mouse CD31 (moCD31, red), and/or human nuclei (HuNu, blue) (e). Quantification of the total length of human (f) and mouse (g) blood vessels. Data are mean ± SD. (isotype IgG: n = 5, ramucirumab: n = 4). Welch’s t test was used to compare with isotype IgG-treated mice. *: p < 0.05. **: p < 0.01. n.s.: not significant.
hypothesize an interaction between the mouse dura mater cells and the human ECs for the development of the human vasculature.

We depict the relationship of engrafted human blood vessels with the original tumor blood vessels in Fig. 3b and Supplementary Fig. S2d. We were unable to identify any factors influencing the engraftment of human blood vessels in the present work. Previous reports indicated that pericytes maintain stability of blood vessels via the angiopoietin 1/Tie2 pathway23 and induce expression of the anti-apoptotic protein Bcl-w in tumor ECs, thereby protecting them from cytotoxic damage25. Therefore, we speculate that engraftment of human blood vessels is dependent on the pericyte coverage of the tumor ECs. In addition, we hypothesize that perfusion of transplanted human blood vessels is dependent on the pericyte coverage of the tumor ECs. To better understand vessel function, it would be necessary to assess vascular permeability, but given the admixture of mouse and human blood vessels in our model, it is difficult to distinguish whether it is the mouse or human blood vessels that are responsible for the permeability. Real-time visualization of vascular permeability is desired and further technical improvement will be required in our model.

We documented perfusion through the human blood vessels in Fig. 3c–e. These results show that the human blood vessels do connect with the murine circulation. Cheng et al. showed that HUVEC-derived human blood vessels connect with mouse blood vessels at the host-implant interface and designated this phenomenon “wrapping-and-tapping” (WAT) anastomosis7. VE-cadherin is an important transmembrane component at the endothelial cell–cell junction. Therefore, human blood vessels in our model may connect to mouse blood vessels via VE-cadherin. It will be intriguing to clarify how cellular junctions between human and mouse ECs are induced by adhesion molecules and the cytoskeleton.

In terms of experimental benefits, our model requires fewer animals, and provides for noninvasive, continuous evaluation of a drug’s effect over time in the same animal, which conventional methods cannot offer. This model therefore allows the study not only of morphological changes such as the length of the human blood vessels but also functional evaluation, such as the degree of perfusion. It is widely accepted that functional normalization of the tumor vasculature is extremely important for improving drug delivery26 and facilitating immune cell access27. Our model allows us to examine the perfusion status of the human blood vessels by using a fluorescence-labeled lectin which specifically binds to human vascular ECs. Our current model would be beneficial for examining the morphological changes of functional human blood vessels exposed to drugs that target tumor blood vessels, which improves the translational potential of pre-clinical animal studies.

An interesting characteristic of our model is that cancer cells cannot be engrafted; however, human blood vessels can be maintained. At present, it is difficult why cancer cells are not engrafted, it is possible that our in vivo model may repulse human cancer cells by unknown immune-like cells. As a tool for screening of drugs affecting human vasculature, our model has benefits. On the other hand, when tumor vasculature needs to be observed in combination with the growth of cancer cells, our model would not be appropriate. To solve this problem, we may need several methodological improvements, e.g. shortening the time required for transportation of the tumor sample from the hospital where patients received surgical resection, using high-grade cancer tissues (i.e. those with greater proliferative potential and resistance to immune attack) or using highly immunodeficient mice, especially NOD-Scid interleukin-2 receptor gamma chain null (NOG/NSG) mice instead of NOD-Scid mice28. Another approach to enable cancer cell growth in our model may be to transplant cancer organoids29 or human mesenchymal stem cells30 together with human tissues including human blood vessels.

Even if cancer cells are engrafted in the cranial window, we would not be able to visualize them in this model. Fonnes et al. showed that injection of near-infrared fluorescently-labeled monoclonal antibody targeting epithelial cell adhesion molecule (EpCAM) as in vivo imaging probe (EpCAM-AF680) to patient-derived tumor-bearing mice enabled visualization of tumor31. HuCD31-AF488 and EpCAM-AF647 injection may serve as an in vivo visualization of interaction of cancer cells and tumor blood vessels.

In summary, we developed a human blood vessel transplantation model, which should provide a useful tool for studying the molecular mechanisms of human blood vessel formation in vivo and for testing drugs for their effects or side effects on human blood vessels in vivo.

**Data availability**

The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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Author contributions
Y.T. and N.T. designed the experiments. Y.T., F.M., Y.H., C.I., H.S. and T.I. performed the experiments. Y.T. and H.S. analyzed the data. C.I., H.K. and N.T. contributed scientific advice. Y.T. and N.T. wrote the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
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