Expanding tubular microvessels on stiff substrates with endothelial cells and pericytes from the same adult tissue

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Abstract

Endothelial cells (ECs) usually form a monolayer on two-dimensional (2D) stiff substrates and a tubular structure with soft hydrogels. The coculture models using ECs and pericytes derived from different adult tissues or pluripotent stem cells cannot mimic tissue-specific microvessels due to vascular heterogeneity. Our study established a method for expanding tubular microvessels on 2D stiff substrates with ECs and pericytes from the same adult tissue. We isolated microvessels from adult rat subcutaneous soft connective tissue and cultured them in the custom-made tubular microvascular growth medium on 2D stiff substrates (TGM2D). TGM2D promoted adult microvessel growth for at least 4 weeks and maintained a tubular morphology, contrary to the EC monolayer in the commercial medium EGM2MV. Transcriptomic analysis showed that TGM2D upregulated angiogenesis and vascular morphogenesis while suppressing oxidation and lipid metabolic pathways. Our method can be applied to other organs for expanding organ-specific microvessels for tissue engineering.

Keywords

Microvessel, endothelial cell, pericyte, culture medium, vascular heterogeneity

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Introduction

Microvessels play essential roles in maintaining normal organ functions. Microvascular dysregulation happens in various diseases, such as fibrosis, tumor, and diabetic microvascularopathy.¹ Microvascular structural stability and proper function rely on microvascular endothelial cells (ECs) and pericytes.² Although many studies have focused on embryonic microvascular development using animal models, in vitro culture models of adult microvessels will provide more insights into the mechanisms of microvascular diseases in adulthood. The isolation and growth of microvessels from adult tissues can provide a platform for research on patient-specific diseases and benefit the development of precision medicine. The culture of adult microvessels will also promote the development of microvascular engineering in vitro.

There have been various in vitro culture models for adult microvessels.³ Active angiogenesis can be induced in

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tissue explant culture models, such as retina and aorta ring explants in a three-dimensional (3D) matrix, however, these models face challenges while being translated into clinical settings. It has been widely used to culture a pure population of primary ECs in most laboratories. The human umbilical vein is a popular source for primary EC isolation. ECs can also be isolated from other organs, for example, the kidney, then purified by flow cytometric sorting. The coculture models of ECs and pericytes have been developed to investigate the mechanisms of pericyte recruitment. The pericytes used in these experiments were usually isolated from the brain, an organ rich in pericytes. In recent years, pluripotent stem cells (PSCs) have been an emerging source for producing ECs, pericytes, and smooth muscle cells. However, it is a challenge for the vascular cells derived from PSCs to reflect the biological characteristics of a specific tissue or organ. Due to the heterogeneity of blood vessels throughout the body, the ideal culture models for adult microvessels would contain microvascular ECs and pericytes from the same adult tissue, which will benefit organ-specific studies.

Materials and methods

Isolation of primary microvessels from adult rats

Male Sprague Dawley (SD) rats aged 8–10 weeks were euthanized by an overdose of isoflurane and bilateral thoracotomy. The subcutaneous soft connective tissue was harvested in an aseptic environment. The tissue pieces were digested in an enzymatic solution containing 2 mg/ml collagenase 1 (Worthington, Cat#LS004196), 5 mg/ml bovine serum albumin (BSA, Sigma, Cat#A1933), and 5 μM Y27632 (Selleck, Cat#S1049) in DMEM (Invitrogen, Cat#12800017).

The cyclic digestion method was used to harvest primary microvessels. Every 10 min, the cell suspension was harvested, a new enzymatic solution was added to the remaining tissue, and the digestion cycle continued until all the tissue pieces were digested. The cell suspension was centrifuged at 1000 rpm for 4 min, and cell pellets were resuspended in phosphate-buffered saline (PBS). All the cell pellets were pooled together and filtered through a 30 μm strainer to remove red blood cells and single cells. The long microvessel segments were collected from the strainer mesh and used for culture.

Culture of primary microvessels

The primary microvessel segments were cultured in regular plastic dishes and plates in different media. EGM2MV medium was purchased from Lonza (Cat# CC-3202). Our custom-made medium was DMEM/F12 (Invitrogen, Cat#12400024), supplemented mainly with 2% fetal bovine serum (FBS, Gibco, Cat#10091148), 100 U/ml penicillin and 100 μg/ml streptomycin (Gibco, Cat#15140122), 5 mM nicotinamide (Sigma, Cat#N0636), 1 mM (N-Acetyl-L-cysteine) NAC (Sigma, Cat#A9165), 50 μM Vitamin C (Sigma, Cat#A4403), 3 μM glutathione (Sigma, Cat#G6013), 10 μg/ml insulin (Aladdin, Cat#I113907), 7.5 μg/ml transferrin (Sigma, Cat#T0665), 40 mM Sodium Selenite (Sigma, Cat#S9133).

TGM2D was supplemented with 2 μM Chir99021 (Sellec, Cat#S2924), 0.2 μM A83-01 (Tocris, Cat#2939), 5 μM Y27632 (Sellec, Cat#S1049), and 10 ng/ml VEGF (Peprotech, Cat#100-20). For the titration of A83-01, the concentrations included 0, 0.1, 0.2, 0.5, 1, and 2 μM.

The cells were cultured in an incubator at 37°C, with 5% CO₂ and 95% humidity. The medium was changed every other day in the first 4 days and every day after that.

Total RNA extraction

Total RNA was extracted from the microvascular cells using Trizol (Invitrogen, Cat#15596026) according to the
manufacture instruction. The cells of a 60 mm dish were homogenized in 1 ml Trizol, centrifuged at 12,000×g for 5 min at 4°C, and then the supernatant was transferred into a tube with 0.3 ml chloroform/isooamyl alcohol (24:1). The mixture was centrifuged at 12,000×g for 10 min at 4°C, the upper aqueous RNA layer was transferred into a new tube with equal volume of isopropyl alcohol, and then centrifuged at 12,000×g for 20 min at 4°C. The RNA pellet was washed with 1 ml 75% ethanol, and then centrifuged at 12,000×g for 3 min at 4°C. The pellet was dried in air for 5–10 min. Finally, 50 μl DEPC water was added to dissolve the RNA, which was qualified and quantified using a Nano Drop and Agilent 2100 bioanalyzer (ThermoFisher).

mRNA library construction

The mRNA was purified by Oligo(dT)-attached magnetic beads and fragmented into small pieces. First-strand cDNA was generated by random hexamer-primed reverse transcription and a second-strand cDNA synthesis. A-Tailing Mix and RNA Index Adapters were added. The cDNA fragments were amplified by PCR, and then purified by Ampure XP Beads and dissolved in EB solution. The product was validated on an Agilent Tech 2100 bioanalyzer. The double stranded PCR products were denatured and circularized by the splint oligo sequence to make the final library, which was amplified with phi29 to make DNA nanoball with more than 300 copies of one molecular, which were loaded into the patterned nanoarray and pair end 150 bases reads were generated on DNBSEQ-T7 platform.

Sequencing analysis

The sequencing data was filtered with SOAPnuke (v1.5.2). The clean reads were matched to the reference genome using HISAT2 (v2.0.4). Bowtie2 (v2.2.5) was applied to align the clean reads to the reference coding gene set. Then expression level of gene was calculated by StringTie (v2.1.2). KEGG and GO analysis were performed in DAVID Bioinformatics Resources (https://david.ncifcrf.gov). The heatmap and bubble plot were drawn by SRplot (http://www.bioinformatics.com.cn) according to the gene expression in different samples.

Quantitative RT-PCR analysis

Total RNA was extracted from the microvascular cells using Trizol (Invitrogen, Cat#15596026) according to the manufacture instruction. RNA quality was checked by measuring the optical density ratio at 260/280 nm on a nanodrop. RNA from each sample was reverse transcribed into cDNA by an Evo M-MLV RT Mix Kit (Accurate Biotechnology, Cat#AG11728). Experiment was carried out using the forward and reverse primers listed below. Fold change in mRNA expression levels was calculated by the comparative Ct method, using the formula 2-(ΔΔCt) and GAPDH as a calibrator. The primer list:

Apn (AAGCCCAAGACTTGAGGAC;GGACGACATATTTC
CGCTTCTG), DiI4 (AGGTGACTCCCGCTGAGC
TGCCGTTAGTGAATTATCC), Kne3 (ACAGATGCGAGA
GTCAGATCG;TGTAGTGTCGGCCGTTG), Pdhd12
(CAGCAGGTCTGAACTGGAGA;TGAGCATCGTGGCTTA
CGGA), Phnx1 (ATGCCCGAGGGCTTCTAGAT;TCTT
CGGTACTCGGGG), Car2, (GCTGGAATGTGTGACC
TGGA;CCACGTCGAGGGTGCTT), Wnt5a (TGGGCA
CATTTCCACGCTG;TGTCCTTGAGAAAGTCCCG), Fzd1
(GCCTCAACACCAGTCCACAA;TGCTTTAAAT
GCCACTCGG), Cxcl12 (AGCCTTAAACAAGAGGCTCA
AG;TGAGGGTGATCTCGCTT), Mmp9 (GATGCC
CAACCTTTACCAG;AAGGTCAAGCCGACCCCTACA).

Immunostaining

The cells were fixed in 4% paraformaldehyde (PFA) for 30 min, washed with PBS for three times, permeabilized by 0.1% Triton for 10 min, blocked in 5% normal donkey serum for an hour, and incubated with primary antibodies in the blocking solution at room temperature. After 2 h of primary antibody incubation, the cells were washed in PBS for three times and incubated with secondary antibodies in blocking solution for 1 h at room temperature. The primary antibodies used in this study included CD31 (Abcam, Cat#ab28364; Santa Cruz, Cat#SC376764; Novus, Cat#AF3628), vWF (Proteintech, Cat#11778-1-AP), LYVE1 (Affinity, Cat# AF4202), NG2 (Proteintech, Cat#55027-1-AP), ZO1 (Proteintech, Cat#21773-1-AP), PDGFRβ (Proteintech, Cat#13449-1-AP), CD45 (Proteintech, Cat#20103-1-AP), and SMA (Santa Cruz, Cat#sc-32251). The secondary antibodies used in this study were purchased from Invitrogen (Cat#A10040, A-21202, A-21447). Cell nuclei were stained with DAPI (4’,6-diamidino-2-phenylindole). Confocal imaging was performed on a Leica SP8 confocal microscope.

Flow cytometry

The primary microvessels were cultured in TGM2D medium for 3 or 7 days before flow cytometric analysis. For flow cytometry, the primary microvessels were dissociated into single cells by accutase (Sigma, Cat#A6964). The staining procedure was performed on ice, including the following: blocking in 3% normal donkey serum for 15 min, incubation in primary antibodies (CD31 from Novus, Cat#AF3628, NG2 from Proteintech, Cat#55027-1-AP, CD45 from Proteintech, Cat#20103-1-AP), secondary antibodies (Invitrogen, Molecular probes, Cat#A10040, A-21202, A-21447). The cells were suspended in the sorting buffer containing 1% BSA and 1 mM EDTA in PBS, went through a single-cell strainer before analysis on a Beckman CytoFLEX LX cytometer.
Statistical analysis

To quantify the features of the microvessels, at least three images randomly selected from each group were analyzed using ImageJ. The total microvessel length and branch point numbers used for analysis were the sums on a 10× image area. Data were presented as means ± SD unless otherwise indicated. One-way or two-way ANOVA was performed on the data of multiple groups, followed by Bonferroni post hoc tests. Student’s t-test was used to analyze the differences between two groups.

Results

Expanding tubular adult microvessels in TGM2D

Primary microvessels were isolated from adult SD rats (Figure 1(a)). The subcutaneous soft connective tissue was harvested from adult rats and digested in an enzymatic solution containing mainly collagenase I at 37°C. We designed a cyclic digestion method to improve the viability of primary microvessels (Figure 1(a)). Every 10 min, we harvested cell suspension from the digestion tube and added a freshly prepared enzymatic solution to the tube. The cell suspension was centrifuged, and the pellets were re-suspended in a collection tube containing phosphate-buffered saline (PBS). The digestion cycle continued until all the tissue pieces were wholly digested. The cell suspensions were pooled together and filtered through a 30 μm strainer to remove red blood cells and other single cells. The microvessel segments were collected from the strainer mesh (Figure 1(a)). We obtained the microvessel segments with a length range of about 50–150 μm (Figure 1(b); Supplemental Figure S1). In the commercially available medium, EGM2MV, ECs formed a monolayer on the rigid surface of plastic culture dishes, consistent with traditional culture methods (Figure 1(c)).

To make a better culture system, we set up to formulate a basal medium that was DMEM/F12 supplemented with some ingredients. Considering that oxidative stress was the primary threat to the cells in vitro, we tested the effect of adding several antioxidants, such as nicotinamide, NAC, glutathione, and Vitamins C, to the DMEM/F12 medium (See Materials and methods). We then screened signaling pathways that may impact the growth of adult microvessels in vitro. Vascular endothelial growth factor (VEGF) was not added in this screening because its potent proangiogenic effect may mask any roles of other signaling pathways. During 2 weeks of culture in vitro, most groups’ culture dishes were covered by lipid-like cells characterized by the accumulation of lipid-like droplets, except those supplemented with Chir99021 (C) (Figure 1(d)). Chir99021 is a GSK3β inhibitor and thus stabilizes β-catenin and activates the canonical Wnt signaling pathway. The result was consistent with a previous report that upregulated Wnt signaling inhibited adipogenesis. Thus, Chir99021 was added to the basal medium for the following study.

ECs may undergo endothelial-to-mesenchymal transition and lose EC identity, during which the TGFβ signaling pathway is a master regulator. We thus examined the effect of an inhibitor of the TGFβ/Smad signaling pathway, A83-01 (A), in combination with Chir99021. We further examined another small molecule, Y27632 (Y), a ROCK inhibitor that was reported to improve embryonic stem cell survival in vitro. The different combinations of these three small molecules were investigated (Figure 1(e)). The primary microvessels were cultured for about a week and analyzed by immunofluorescence staining. We found that the combination of the three small molecules, C + A + Y, significantly promoted the growth of tubular microvessels (Figure 1(e) and (f)). The addition of VEGF further significantly increased microvessel length and branch point number (Figure 1(e) and (f)). We referred to the medium containing C, A, Y, and VEGF as TGM2D, which significantly promoted the growth of adult tubular microvessels in vitro, compared to the commercially available medium EGM2MV (Figure 1(g)).

TGM2D upregulated angiogenesis and downregulated oxidation and lipid metabolism

We performed mRNA sequencing to analyze gene expression profiles of the microvessels in TGM2D (T) and EGM2MV (E) compared to fresh primary microvessels (V). We focused on the genes up- or down-regulated in EGM2MV and TGM2D (Supplemental Figure S2, A). KEGG pathway analysis showed that EGM2MV specifically upregulated the pathways of oxidation and lipid metabolism (Supplemental Figure S2, B). GO analysis showed that the elevated biological processes in EGM2MV included mainly oxidation, lipid metabolism, and aging, which involved increased mitochondrial activities (Figure 2(a); Supplemental Figure S2, C). On the contrary, oxidation- and lipid metabolism-related processes were significantly suppressed in TGM2D (Supplemental Figure S2, A). TGM2D specifically upregulated Notch, TNF, Rap1, Ras, PI3K-Akt, and chemokine signaling pathways, which were closely related to angiogenesis (Supplemental Figure S2, B). GO analysis showed that the upregulated biological processes in TGM2D included mainly angiogenesis, vascular branch formation, vascular remodeling, EC proliferation, and migration, which mainly involved cell-cell junctions between ECs, contributing to the structural integrity of microvessels (Figure 2(b); Supplemental Figure S2, D).

The genes for the identity of ECs (Cd31, Cdhs5, Tie115,16) and endothelial progenitor cells (Cd34, Kdr, and Kit) were down-regulated in EGM2MV but significantly upregulated in TGM2D (Figure 2(c)). Sox family members, including Sox7, 17, and 18, were also upregulated in
TGM2D for the differentiation and maturation of ECs.\(^{18}\) TGM2D also upregulated the expression of Apln, which was recently reported to mark an EC subpopulation contributing to vascular development\(^{19,20}\) and regeneration.\(^{21,22}\) Proangiogenic genes were significantly upregulated in TGM2D, such as Mmp9,\(^{23}\) Angpt2,\(^{24}\) Wnt signaling genes (Wnt4, Wnt5a, Cpz, Sfrp2),\(^{25}\) and Notch signaling genes (Notch1, Notch3, Dll4, Jag1, Hey1),\(^{26-28}\) and chemokine signaling (Cxcl12, Cxcr4, and Ackr3)\(^{29,30}\) (Figure 2(c) and (d)). Genes significantly upregulated in TGM2D also included: Kcne3,\(^{31}\) Esm1,\(^{32}\) as tip EC markers; Car2, which promoted EC survival,\(^{33}\) Pcdh12, a protocadherin,\(^{34}\) Plxnd1, for EC mechanotransduction.\(^{35}\)

In addition to the markers of ECs, TGM2D also upregulated Pdgfb and Pdgfrb, which were expressed in ECs and pericytes respectively for pericyte recruitment.\(^{36}\) Other pericyte markers, including Mcam (Cd146) and Cspg4 (Ng2)\(^{37}\) were also upregulated in TGM2D. These results indicated that both ECs and pericytes of the same microves sel segments were expanded in TGM2D.
Individual roles of Chir99021, Y27632, and A83-01 in tubular microvessel growth in vitro

To investigate the roles of the small molecules in TGM2D, VEGF was supplemented to the basal medium to obtain enough microvessels for analysis. The primary microvessels were cultured in different groups of the combinations of the three small molecules. During a week of culture in vitro, a single small molecule Chir99021 alone was enough to promote tubular microvessels’ growth significantly. Adding Y27632 further increased microvessel length and branch point number. Adding A83-01 to C or CY groups did not
benefit microvessel length or branch point number but produced some local areas with wide EC bundles (Figure 3(a) and (b)).

Transcriptomic analysis showed that the upregulated expression of most proangiogenic genes was mainly attributed to Chir99021, which acted as a Wnt agonist and induced the activation of the Notch signaling and other pathways associated with angiogenesis, branch formation, and cell migration (Figure 3(c) and (e)). A83-01 played a minor role in angiogenesis but contributed to preserved EC functions by upregulating the expression of genes related to cholesterol efflux (Apoe), anti-coagulation (Thbd), and cell proliferation (such as Pgf, Igf1, and Adm.) (Figure 3(d) and (f)). Both Chir99021 and A83-01 promoted the expression of the progenitor marker CD34, indicating that EC progenitor properties were maintained and EndoMT was suppressed (Figure 3(d)). Interestingly, the addition of Y27632 did not significantly change the gene expression profiles of the microvessels (data not shown), although there was indeed an improvement in the microvascular length and branch formation (Figure 3(a) and (b)). Y27632 may only take effect at the protein level through regulating cell contractility.

**Figure 3.** The roles of individual small molecules: (a) immunofluorescence images of the microvessels in different groups. The antibody against CD31 was used to label endothelial cells. DAPI was used to label cell nuclei. Scale bars, 100 μm. (b) quantification of microvessel length and branch point number in different groups (n = 6). Data were presented as mean ± SD. One-way ANOVA was performed on the data, followed by Bonferroni post hoc tests. *p < 0.05, **p < 0.01. (c and d) the heatmaps of genes upregulated by Chir99021 (c) and A83-01 (d) in different groups, and (e and f) GO analysis of the biological processes upregulated by Chir99021 (e) and A83-01 (f).
**Long-term culture of adult tubular microvessels in vitro**

We next examined the long-term performance of the three small molecules in the growth of microvessels in vitro. For a total of 8 weeks of culture, the cells were fixed and immunostained for EC marker CD31 at different time points: 1, 2, 3, 4, and 8 weeks. Although Chir99021 significantly promoted the growth of microvessels during the first week (Figure 3), the microvessels in the Chir99021 medium degraded quickly from the second week (Figure 4(a), (e), and (f)). The addition of A83-01 to Chir99021 medium slightly rescued the microvessels from the second week (Figure 4(b), (e), and (f)). Y27632 had a noticeable effect on the survival of microvessels from the second week (Figure 4(c)–(f)). The full TGM2D, containing C, A, and Y, could sustain microvascular growth for 8 weeks (Figure 4(d)–(f)). Microvascular length and branch formation were relatively high for at least 4 weeks (Figure 4(d)–(f)). The wide EC bundles in the groups of CA and CAY disappeared, and all the ECs adopted a tubular structure from the second week, indicating that active microvascular remodeling happened during culture in vitro (Figure 4(b) and (d)).

**Preserving syngeneic pericytes around tubular microvessels in vitro**

There were few LYVE1+ lymphatic endothelial cells in the culture, indicating that our custom-made TGM2D could specifically expand tubular blood vessels (Figure 5(a)). The adult tubular microvessels also showed strong endothelial marker vWF (Figure 5(b)) and the tight junction protein ZO1 (Figure 5(c)). More importantly, plenty of pericytes around the microvessels expressed the markers including NG2, PDGFRβ, and αSMA (Figure 5(d)–(f)).

The digestion procedure would give rise to microvessels and other types of cells, including adventitial cells, fibroblasts, and immune cells. Our method excluded most of these single cells by cell straining and obtained a relatively pure population of microvessel segments from the strainer mesh (Figure 1(b)). We performed flow cytometry and immunostaining analysis to examine whether there were immune cells in the culture (Figure 6).

We tried digesting the primary microvessels into single cells by incubating them in the enzymatic solution for a longer time, which led to cell death. In our hands, it was not easy to dissociate the primary microvessels into single cells during the first 2 days of culture. Thus, we performed flow cytometric and immunostaining analysis on the primary microvessels at 3 and 7 days. The results showed very few CD45+ cells in the microvascular culture (Figure 6(a) and (b)). There were about 10% of CD31+ ECs and 9% of NG2+ pericytes on day 3, which were around 4% and 20%, respectively, on day 7 (Figure 6(c)). By immunostaining, we also found many NG2+ pericytes (Figure 6(d), arrow) and a few CD31+ ECs (Figure 6(d), arrowhead) expressing the proliferation marker Ki67 on day 7. These results suggest that pericytes actively proliferated in vitro, which may explain the lower percentage of ECs during culture. We also noticed some CD31−NG2−CD45− cells in the areas far from microvessels, which may be derived from pericyte differentiation or residual fibroblasts. We will evaluate the significance of these uncharacterized cells and investigate them in our future studies.

**The balance between the ECs and pericytes**

TGFβ signaling pathway is essential for the differentiation and maturation of vascular mural cells, including smooth muscle cells and pericytes. There were still some NG2+ pericytes around microvessels in a week of culture in TGM2D containing A83-01 (Figure 5). Thus, we investigated whether NG2+ pericytes still existed in a long-term culture of 2 weeks, when TGFβ signaling was inhibited by different concentrations of A83-01. We found that higher concentrations of A83-01 induced a higher density of microvessels and the concentration of 0.5μM gave the maximal level of microvessel length and branch formation (Figure 7), which was consistent with previous report. However, there were few NG2+ pericytes when A83-01 concentration was bigger than 0.2μM. (Figure 7). Restoration of the TGFβ signaling by removing A83-01 from the medium at the second week rescued NG2+ pericytes to some extent (Figure 8). However, NG2+ pericytes cannot be recovered if A83-01 was removed at the third week (data not shown).

**Discussion**

This study presented a model platform for expanding tubular microvessels with the ECs and pericytes from the same tissue on the rigid surface of regular culture dishes. The subcutaneous soft connective tissue of the adult body contains microvessels, fibroblasts, and some stem cells. It can be relatively easily obtained during minor surgeries, such as liposuction, plastic surgeries, and foreskin surgeries, making it a promising source for translational medicine. In traditional EC isolation methods, the tissues were digested for a long time until single ECs were obtained. Long-term exposure to digestive enzymes may harm the cells to some extent. The isolated single ECs lost their microenvironment, including cell-cell connections between ECs and between ECs and pericytes. In this study, we developed a cyclic digestion method to maximize the viability of primary cells and the structural integrity of microvessels. This method can be applied to primary cell isolation from other tissues and organs in general.
Figure 4. Time-lapse analysis of the microvessels in different media: (a–d) immunofluorescence images of the microvessels in the media supplemented with C (a) CA (b), CY (c), and CAY (d) in the presence of VEGF. The antibody against CD31 was used to label endothelial cells. DAPI was used to label cell nuclei. Scale bars, 100 μm. (e and f) quantification of microvessel length and branch point number in different groups at different time points (n = 6). Data were presented as mean ± SD. Two-way ANOVA was performed on the data, followed by Bonferroni post hoc tests. ***p < 0.001.

The maintenance of tubular morphology of adult microvessels in this study was achieved through the orchestration of multiple pathways. One of the main threats to the cells cultured in vitro is oxidative stress, which induces DNA damage, senescence, and apoptosis. Oxidation was also a major biological process in the culture of EGM2MV (Figure 2(a)). The addition of antioxidants may contribute to the downregulation of oxidation pathways in TGM2D. The
Figure 5. Endothelial and pericyte marker expression in TGM2D. The microvessels were cultured in TGM2D for 1 week and then immunostained by the antibodies against LYVE1 (a), vWF (b), CD31 (c–f), ZO1 (c), NG2 (d), PDGFRβ (e), and αSMA (f). Nuclei were stained by DAPI. Scale bars, 100 μm.
GSK3β inhibitor Chir99021 upregulated Wnt signaling and played a dual role in the system. Activated Wnt signaling has been reported to suppress adipogenesis. On the other hand, Wnt signaling was a master regulator in angiogenesis, blood vessel morphogenesis during embryonic development and adult tissue regeneration. Our results showed that the treatment of Chir99021 alone could activate most of the proangiogenic genes of adult microvessels (Figure 3(c) and (e)), indicating that Wnt signaling also played a central role in the tubular microvessel growth in vitro.
TGFβ signaling pathway is essential for vascular development as it modulates the differentiation and maturation of vascular mural cells, including smooth muscle cells and pericytes. On the other hand, TGFβ signaling can induce the loss of EC identity through endothelial-to-mesenchymal transition (EndoMT). TGFβ inhibition maintained EC identity and proliferation. Our results showed that the inhibition of TGFβ/Smad signaling by A83-01 upregulated the expression of Cd34, a marker for endothelial progenitor cells. Treatment of A83-01 also promoted the expression of the genes for EC proliferation, such as Pgf, Igf1, and Adm. More importantly, the critical functional genes, such as Apoe for cholesterol transport and Thbd for anti-coagulation, were upregulated by the treatment of A83-01 (Figure 3). All these results indicate that the inhibition of TGFβ/Smad signaling by A83-01 promoted EC proliferation and maintained EC functions.

Y27632 is a ROCK inhibitor that has been widely used in the culture of pluripotent stem cells and functions to promote cell survival in vitro. Studies in corneal endothelium showed that Y27632 promoted cell proliferation by upregulating Cyclin D and downregulating p27 gene expression. However, few genes were influenced by the addition of Y27632 in our study (data not shown), indicating that...
Y27632 may take effect at the protein level. Y27632 has been reported to reduce cell contractility by down-regulating the phosphorylation of non-muscle myosin light chain II (MLC). As pericytes wrap around microvessels and exert contractile forces to the EC tubules and limit their growth, Y27632 may promote microvascular tubule growth and branch formation by reducing the contractility of ECs and pericytes, which needs further investigation.

**Figure 8.** Dynamic regulation of NG2 pericytes by A83-01: (a–c) the primary microvessels were cultured in CY for 14 days (a), in CAY for 14 days (b), and in CAY for 7 days, followed by CY for another 7 days (c), then immunostained by antibodies against CD31 and NG2. Nuclei were stained by DAPI. Scale bars, 100 μm. (d) quantification of microvessel length, NG2⁺ pericyte number per microvessel length, and NG2⁺ pericyte coverage ratio of total microvessel length (%). Data were presented as mean ± SD. Student’s t-test was performed to analyze significant differences between groups. ***p < 0.001.
There have been several models for engineering tubular microvessels in vitro. A classic method is to culture ECs on the surface of Matrigel, and ECs will self-assemble into tubules and form a network, which, however, can only survive a few days.\textsuperscript{3} The ECs can also self-assemble into tubular structures inside a 3D matrix, such as collagen, Matrigel, and fibrin gels.\textsuperscript{3} Microfluidics techniques can make these 3D microvessels perfused, better-mimicking microvessels in vivo.\textsuperscript{3} These tubular microvessels in 3D hydrogels were usually maintained and investigated for up to 2 weeks.\textsuperscript{3}

Most microvascular diseases are chronic diseases and take place in years or decades. The potential effect of a treatment may not emerge in a short time. A goal in the research of microvascular engineering is to maintain the structure of microvessels for a longer time. In this study, the microvessels in the experimental groups of C, CA, CY, and CAY had minor differences at the first week time point, but they behaved distinctly from the second to eighth week (Figure 4). The microvessels in group C quickly deteriorated in the second week, indicating that the activated Wnt signaling pathway alone promoted angiogenesis but was not enough to maintain the structural stability of microvessels in vitro. The beneficial effect of Y started to emerge from the second week. Adding Y to the media C or CA significantly improved microvessel length and branch formation. The addition of A to CY medium further promoted microvessel stability, possibly through upregulating the genes related to EC functions and proliferation (Figure 3). The microvessel length and branch formation in TGM2D (VEGF + CAY) were maintained at a relatively high level for at least 4 weeks in vitro (Figure 4). The time period of the third and fourth weeks had relative stable microvessels, which are a proper candidate for experimental manipulations (Figure 4).

There was a subtle balance between the ECs and pericytes in this model. Pericytes play a vital role in regulating microvessel structural stability and functions.\textsuperscript{36} NG2 is a glycoprotein located on the plasma membrane of pericytes and regulates microvessel integrity.\textsuperscript{36} Genetic knock-out studies showed that NG2\textsuperscript{−/−} animals had fewer microvessels than normal controls.\textsuperscript{36} Our results are consistent with previous reports that TGFβ signaling is essential for the differentiation and maturation of pericytes.\textsuperscript{38} Although there were slight differences in NG2\textsuperscript{+} pericytes between CY and CAY media at the first week, a long-term culture of 2 weeks in CAY medium significantly reduced the number of NG2\textsuperscript{+} pericytes around microvessels (Figures 7 and 8). NG2\textsuperscript{+} pericytes can be preserved by removing A83-01 from the medium from the second week (Figure 8), but they will vanish if A83-01 was removed from the third week. These results indicate that the maintenance of NG2\textsuperscript{+} pericytes requires TGFβ signaling. On the other hand, TGFβ signaling played an opposite role in ECs. TGFβ inhibition was required to maintain EC identity and functions in vitro.\textsuperscript{13,39} Future studies are warranted to investigate how to coordinate the growth of syngeneic ECs and pericytes in tubular microvessels in vitro.

Our method will have broad applications in microvascular engineering and regeneration. It can be used in tissue vascularization, which is a core question in tissue engineering.\textsuperscript{57} For example, intensive efforts have been made to fabricate vascularized tissue and organ models, such as the vascularized pancreas,\textsuperscript{58,59} adipose tissue,\textsuperscript{60} bladder,\textsuperscript{61} islet,\textsuperscript{62} skin,\textsuperscript{63} and so on. The model presented in this study can be further improved and applied in these ex vivo settings in the future. One critical aspect is perfusion, which is essential for the study of microvascular functions, including permeability,\textsuperscript{64,65} mechanobiology,\textsuperscript{66} tumor cell and immune cell infiltration.\textsuperscript{67} More advanced microvascular models can be made based on our method in combination with other engineering technologies, such as microfluidics,\textsuperscript{68} electrospinning,\textsuperscript{69} microspheres,\textsuperscript{68} and 3D printing.\textsuperscript{57}

Conclusion

Our study established a method for expanding tubular microvessels on stiff substrates of regular culture dishes and retaining the ECs and pericytes from the same adult tissue. This model will hopefully provide more insights into the mechanisms of EC-pericyte interactions in tissue regeneration and diseases. The method established in this study also applies to other organs for engineering organ-specific microvascular models. As human microvessels can be relatively easily obtained from the subcutaneous soft connective tissues during minor surgeries, the platform presented in this study can hopefully be translated to clinical settings, benefitting the development of personalized microvascular engineering for precision medicine.

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Author contributions

DW designed the experiments. DW and JL wrote the manuscript. XS, YY, YL, YX, and DW performed primary cell isolation, cell culture, immunostaining and confocal microscopy. JM performed qRT-PCR. YL and LM performed flow cytometric analysis. ZW performed quantification of microvessels. HZ, XQ, PL, XS, YY, YL, YX, and DW performed primary cell isolation, cell culture, immunostaining and confocal microscopy. JM performed qRT-PCR. YL and LM performed flow cytometric analysis. ZW performed quantification of microvessels. HZ, XQ, PL, XS, YY, YL, YX, and DW performed transcriptomic analysis.

Availability of data and materials

The data required to reproduce these findings can be made available upon reasonable request.
Declaration of conflicting interests
The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Ethical approval
The animal procedure was conducted according to the Ministry of Science and Technology guide for laboratory animal care and use and approved by the animal care and use committee of Qingdao University.

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