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In Vitro and In Vivo Evaluation of 3D Bioprinted Small-diameter Vasculature with Smooth Muscle and Endothelium

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Abstract

The ability to fabricate perfusable, small-diameter vasculature is a foundational step toward generating human tissues/organs for clinical applications. Currently, it is highly challenging to generate vasculature integrated with smooth muscle and endothelium that replicates the

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Author contributions

H. C. and L. G. Z. conceived the ideas and designed the experiments. H. C., W. Z., Y. C., S. L., and X. Z. conducted the experiments. H. C., M. N., and S. M. performed data analysis. Y. H., C. L., Z. Y., Y. Z., S. W., M. M., and K. H. designed and carried out animal experiments. H. C., M. N., S. M., and L. G. Z. prepared the manuscript.

Appendix A. Supplementary data

Supplementary data to this article can be found online at

Competing financial interests

The authors declare no competing financial interests.

Data availability

The authors declare that all data supporting the findings of this study are available within the paper and its supplementary information, source data for the figures in this study are available from the authors up on request.

complexity and functionality of natural vessels. Here, a novel method for directly printing self-standing, small-diameter vasculature with smooth muscle and endothelium is presented through combining tailored mussel-inspired bioink and unique “fugitive-migration” tactics, and its effectiveness and advantages over other methods (i.e., traditional alginate/calcium hydrogel, post-perfusion of endothelial cells) are demonstrated. The biologically inspired, catechol-functionalized, gelatin methacrylate (GelMA/C) undergoes rapid oxidative crosslinking *in situ* to form an elastic hydrogel, which can be engineered with controllable mechanical strength, high cell/tissue adhesion, and excellent bio-functionalization. The results demonstrate the bioprinted vascular construct possessed numerous favorable, biomimetic characteristics such as proper biomechanics, higher tissue affinity, vascularized tissue manufacturing ability, beneficial perfusability and permeability, excellent vasoactivity, and *in vivo* autonomous connection (~2 weeks) as well as vascular remodeling (~6 weeks). The advanced achievements in creating biomimetic, functional vasculature illustrate significant potential toward generating a complicated vascularized tissue/organ for clinical transplantation.

Keywords

3D bioprinting; mussel-inspired; small diameter vasculature; endothelium; smooth muscle

1. Introduction

The crux of engineering tissues/organs in regenerative medicine is to maintain metabolic functions of cellularized tissue scaffolds *in vivo*. Vascular regeneration is a necessary prerequisite for the fabrication of functional tissues/organs that recapitulate the multi-scale structural, mechanical, and physiochemical aspects of biological functions for future clinical tissue/organ implantation applications [1–5]. Large-diameter vascular grafts (diameter >5 mm) are commercially available for clinical use; the capillaries, however, only consist of endothelial cells (several micrometers to tens of micrometers thick), which can be spontaneously vascularized through sprout angiogenesis *in vivo* [2, 5, 6]. On the contrary, manufacturing small-diameter vasculature, bridging large arteries/veins and capillaries, remains a huge challenge for this field [7, 8].

Many attempts to fabricate small-diameter vasculature have been made in tissue engineering [9–12], including various 3D bioprinting techniques [5, 13–17], to mimic a vascular network that can support medium flow and oxygen supplementation for cell viability, promoting improved tissue maturation and development. Although impressive successes have been achieved, these studies have only generated endothelium via colonizing endothelial cells in the perfusable channel [5, 18]; the native structure is much more complicated. Native muscular small-diameter arteries or veins have two or three distinct tunics [11, 19]. The intima, the innermost layer, is a thromboresistant confluent monolayer of endothelial cells; the media, the middle layer, is comprised of a dense population of concentrically organized smooth muscle cells with bands or fibers of elastic tissues; and the adventitia, the outermost layer, is a collagenous extracellular matrix (ECM) containing mainly fibroblasts and perivascular nerves [11, 19]. As functional vasculature, from a pathophysiological and clinical perspective, both endothelial cells and smooth muscle cells play central roles in

regulating hemodynamic forces, maintaining homeostasis, and facilitating remodeling [20–22]. Endothelium-lined vessels play an important role in maintaining the vessel wall permeability barrier, regulating coagulation (thromboresistant), and controlling blood flow, while the smooth muscle layer is vital in generating contractile forces of vasoreactivity, allowing a tolerance of systemic arterial pressures, regulating hemodynamic behaviors and other physiological activities [19, 23, 24]. Without modulation of the vascular smooth muscle, the vascular geometries present at branch points, curvatures, and post-stenotic regions would increase flow pattern disturbances, leading to endothelial dysfunction and other pathophysiological consequences, such as atherosclerosis [25, 26].

Currently, generating the stratified and self-standing vessels that replicate the complexity and functionality of native vasculature, facilitating the *in vivo* remodeling process and connection, remains a significant challenge. Given these requirements, it is envisaged that biomaterials play an important role in successfully engineering biomimetic vasculature. It should possess several specific characteristics: rapid and precise modeling ability; structural stability for long-term implantation; proper mechanical elasticity; excellent cell viability, adhesion, and proliferation; beneficial perfusability and permeability; biodegradability; biocompatibility; as well as bioactivity.

In this study, coaxial extrusion printing of self-standing, small-diameter vasculature with stratified architecture was explored using tailored bioinks. The biologically inspired catechol-functionalized gelatin methacrylate (GelMA/C), was designed and synthesized to provide the spectrum of chemical (in situ controllable crosslinking), biological (bio-functionalization), and physical (interstitial flow and elastic properties) cues for vascular bioprinting. Additionally, a fugitive crosslinking slurry was cooperatively used to support and solidify the GelMA/C bioink, thereby ensuring the successful generation of the vasculature. The stratified, independent, and free-standing architecture facilitates its integration into any engineered tissue implants, compared with dependent variations [8, 27]. To fully exploit the benefits of the 3D bioprinted vasculature for tissue regeneration, a systematic investigation both *in vitro* and *in vivo* was performed, further verifying the efficiency and advanced potential for scaling up for complex tissue/organ applications.

2. Materials and methods

2.1 Materials.

All chemical reagents were from Sigma-Aldrich and were used as received unless otherwise stated. All materials in the experiments involving cell culture and animal study are pre-processed using standard sterilization methods.

2.2 Synthesis of reactive gelatin methacrylate/catechol (GelMA/C).

GelMA/C was synthesized through a two-step chemical reaction, in which gelatin (type A; 300 bloom from porcine skin) was successively reacted with methacrylic anhydride and dopamine. Briefly, 1 g Gelatin (Gel) was dissolved in 10 mL 0.01 M phosphate buffered saline (PBS) at 60 °C. Methacrylic anhydride (10 mL) was slowly added into the 10% gelatin solution while stirring, and then reacted at 60 °C for 3 h. The product was dialyzed

against deionized water for 3 d. The dialyzed solution was lyophilized to obtain pure GelMA. 1 g GelMA was dissolved in 20 mL degassed mixed solvent (2 x PBS (pH=5.0) and dimethylformamide (DMF), v/v=1/1) at 60 °C. 1-ethyl-3-(3-(dimethylamino)-propyl) carbodiimide (EDC) and N-hydroxysulfosuccinimide (NHS) were added to the solution with excess molar ratio (carboxyl of GelMA/EDC/NHS=1:5:5). After activation for 1 d, dopamine hydrochloride was added to the solution with different molar ratios to carboxyl and stirred overnight. The crude product was purified by dialysis for 3 d in deionized water (pH=5) and was subsequently lyophilized, which finally resulted in a white or silver GelMA/C with an average yield of 85%. The graft ratio of catechol groups in samples was calculated by measuring absorbance at 280 nm using a UV-visible spectrophotometer. The quantitative measurement was performed using a dopamine standard. The GelMA/C product was characterized using Hydrogen-1 Nuclear Magnetic Resonance (¹H NMR), and its molecular weight was measured by gel permeation chromatography (GPC).

2.3 Formation of GelMA/C hydrogels and characteristics.

The GelMA/C was dissolved in PBS and then adjusted to a neutral pH of approximately 7.8 using 1 M sodium hydroxide (NaOH). To form the GelMA/C hydrogel, the GelMA/C solution was mixed with a sodium periodate (NaIO₄) solution at an equal molar ratio to catechol groups. To determine the swelling properties of the hydrogels, they were incubated in Dulbecco's modified Eagle's medium (DMEM) at 37 °C. The wet weight of the hydrogel constructs was measured at several time points during incubation. The swelling ratio of the hydrogel was calculated using the following equation:

$$SR = \frac{W_s - W_i}{W_i} \times 100\%$$

(Equation. 1),

W_s indicates the weight of the swollen hydrogel at each time point, and W_i indicates the initial weight of the original hydrogel on day 0.

Oscillatory rheology was performed with a Discovery HR-2 rheometer (TA Instruments). The GelMA/C solution was added directly onto the plate. After mixing with the crosslinker, a brown gel plate was immediately prepared with a diameter of 20 mm and an average thickness of 0.5 mm. The strain sweep was performed from 0.01% to 20% at a constant frequency of 1 rad/s, and a frequency sweep was performed from 0.01 to 100 rad/s at a constant strain of 1% in the linear elastic region. No pre-stress was applied to the gels for these measurements.

The adhesion strength of the hydrogel was measured by recording detachment stress of GelMA/C hydrogel from the tissue [28, 29]. Briefly, two pieces of thin muscle tissue were clamped to each side of the tensile tester. A pre-gel solution of GelMA/C (200 μL) was placed between the tissues and then crosslinked by adding oxidant for 10 min. The cohesive sample was pulled with a crosshead speed of 1 mm/min until detachment.

2.4 Cell culture and maintenance.

Human Coronary Artery Smooth Muscle Cells (HCASMCs, ATCC, The Global Bioresource Center) were cultured in smooth muscle growth medium for cell proliferation consisting of Medium 231 supplemented with Smooth Muscle Growth Supplement (SMGS). For smooth muscle differentiation studies, the Smooth Muscle Growth Supplement (SMGS) was replaced by the Smooth Muscle Differentiation Supplement (SMDS). Human Umbilical Vein Endothelial Cells (HUVECs, ThermoFisher Scientific) were cultured in endothelial growth medium (EGM) consisting of Medium 200 and low serum growth supplement (LSGS). Human bone marrow-derived mesenchymal stem cells (hMSCs, Texas A&M Health Science Center, Institute for Regenerative Medicine) were cultured in mesenchymal stem cell growth medium (MSCGM) consisting of alpha minimum essential medium, 20% fetal bovine serum (FBS), 1% L-glutamine, and 1% penicillin/streptomycin. For the following co-culture study, vascular constructs will be optimally maintained in the mixed medium with 1:1, endothelial growth medium: SMC growth medium (or SMC differentiation medium), for further characterization and *in vitro* cell studies. All experiments were performed with HCASMCs, HUVECs, and hMSCs of six cell passages or less.

2.5 3D bioprinting of cell-laden vascular constructs.

Vascular constructs were created on a customized multi-material 3D bioprinter with a coaxial needle extrusion system designed and assembled in our lab. This bioprinter consists of a transparent glass platform, equipped with a LED light array on the bottom, that controls the Z-axis motion, and an X-Y tool head coupled with coaxial nozzle connecting to two separate syringes for feeding two different bioinks simultaneously. Each ink is reserved in a syringe and deposited by applying mechanical pressure. All printing configurations are controlled by the MatterControl 1.5 software package. The printing parameters, such as printing speed (0 ~ 100 mm/s) and extrusion flow rate (0 ~ 1.0 mL/s), were varied in order to obtain the best printing resolution. The coaxial needle gauges were also varied to fabricate various sizes of vessel. The inner diameter (ID) can be adjusted in the range from 500 to 1500 μm using different coaxial needles, and the wall thickness (WT) can be controlled in the range from 100 to 300 μm . Here, tubular lattices (interconnected vascular channels) were printed at room temperature under a sterile environment with a 5-mm line distance, and stacked layer by layer to form a cylindrical structure, mimicking vascular models within living tissue. A bioink mixture containing 20 wt% GelMA/C, and HCASMCs (1×10^6 cells/mL) was extruded through an external needle (18G, diameter: 840 μm) while a crosslinking slurry containing Pluronic F127 (30 wt%), sodium periodate (NaIO_4 , 23.4 nM), and HUVECs (1×10^7 cells/mL) flowed through an internal needle (22G, diameter: 406 μm). Upon printing, chemical gelation was achieved by diffusion of NaIO_4 from the encapsulated slurry. Next, the 3D printed vascular construct was cultured in the cell medium and was flipped every 2~4 hours for up to 24 hours. Most HUVECs were floated in the hollow channel for the following cell attachment. An open, self-standing blood vessel with a bilayered cell structure was formed after removing the fugitive slurry.

Photographs and videos of vascular bioprinting were acquired using a camera and processed by imaging software. Fluorescence dyes were used to improve visualization of GelMA/C ink

(Fluorescein) and crosslinking slurry (Nile red). Morphology of the 3D printed vascular constructs was observed by optical microscopy (Mu800, Amscope) with a video camera. The 3D optical map images were also performed using microscopic CCD accessories. The lyophilized morphology and microporous structure of 3D printed tubular constructs were studied with a scanning electron microscope (SEM) (FEI Teneo LV FEG). All scaffolds were coated with a 10 nm thick gold layer and imaged using a 5-kV electron beam. To visually assess cell location or arrangement in 3D printed vasculature, a fluorescent-labeled cell study was conducted. Before cell printing, HCASMCs, and HUVECs were incubated with CMFDA, and CMTDX (10 μ M Molecular Probes, CellTracker™ Dye, life technologies) each for 30 min at 37 °C, respectively. After 3 days of culture, the constructs were imaged with a confocal microscope.

2.6 Cell viability and function assay.

To distinguish cell viability and proliferation after bioprinting, HUVECs and HCASMCs (1×10^5 cells/mL) were separately printed into the different vascular constructs. A Live-Dead assay kit (BioVision) was performed to evaluate cell viability. Printed cell-laden vascular constructs were placed onto a glass substrate and then treated with calcein-AM (2 μ M) and propidium iodide (4 μ M) for 20 min. The samples were fixed in a GelMA hydrogel, and then performed a cross-sectional cut. Cross-sectioned samples were observed and imaged using a Zeiss 710 confocal microscope. Additionally, cell activity in the printed constructs was quantitatively investigated using a CCK-8 kit. After the predetermined period, the incubation medium was changed with WST-8 solution (10% v/v in medium) (Dojindo). After 4 h of incubation, the absorbance values of the supernatant solution were measured at 450 nm on a photometric plate reader (Thermo Scientific).

Immunostaining was used to assess the cellular bioactivity for both HCASMCs and HUVECs in the 3D vasculature. The printed constructs were first fixed in 10% buffered formalin for 20 min. After fixing, constructs were permeabilized in 0.1% Triton X-100 for 30 min, and then blocked using 2% bovine serum albumin (BSA) for 2 hours. Primary antibodies were incubated overnight with the constructs at 4 °C, and then secondary antibodies were incubated with the constructs for 2 hours. The cells' cytoskeleton was identified with double staining of F-actin (red) using Texas Red-labeled phalloidin for 30 min and nuclei (blue) using 4, 6-diamidino-2-phenylindole dihydrochloride (DAPI) (Invitrogen) for 10 min. Perfused washing with PBS was required for each step. Samples were observed and imaged using a confocal microscope. The following primary antibodies were used for staining: Rabbit monoclonal anti-alpha smooth muscle actin (α -SMA) antibodies (Abcam) and goat polyclonal anti-von Willebrand factor (vWF) antibodies (Santa Cruz Biotechnology). Goat anti-rabbit IgG H&L Alexa Fluor® 488 (Abcam) and chicken anti-mouse IgG-TR (Santa Cruz Biotechnology) were used as secondary antibodies in current study.

2.7 3D bioprinting of vascularized tissue model.

For vascularized tissue/organ printing, the zoning fabrication was employed through a dual bioprinting manufacturing technology. Primary cell-laden matrices with designed architectures were fabricated using a stereolithography apparatus (SLA) based 3D bioprinter

(the laser beam is about 190 μm , the wavelength is about 355 nm, and intensity output of emitted UV is $\sim 20 \mu\text{J}$ at 15 kHz), while the vasculature was printed by coaxial extrusion printer as mentioned in Section 2.5. In a typical experiment, a prepolymer mixture in PBS containing 10 wt% GelMA, photoinitiator (Irgacure 2959, 0.5 wt%) and mesenchymal stem cells (MSCs) was prepared as the photocurable bioink for 3D tissue matrix bioprinting. MSC-laden tissue matrix was printed by SLA bioprinter, in which the printed vasculature was embedded during the SLA bioprinting. To visually assess cell location or arrangement in 3D printed vascularized tissue, a fluorescent-labeled cell study was conducted. HCASMCs, HUVECs, and hMSCs were incubated with CMFDA, CMTDX and CMTMR (10 μM Molecular Probes, CellTracker™ Dye, life technologies) each, respectively, for 30 min at 37 °C before cell printing. After 3 days of culture, the constructs were imaged with a confocal microscope. Similar to the method above, a Live-Dead assay was also performed to evaluate MSC viability in the 3D vascularized tissue model when perfusing medium through 3D bioprinted vasculature. The medium was perfused into the channel at a flow rate of 6.0 mL/min for 24 h.

2.8 Comparative study of alginate/calcium bioink system.

The combination bioinks, composed of alginate (6 mg/mL) and calcium chloride (CaCl_2 , 6 mg/mL) were used to fabricate the vasculature by coaxial extrusion bioprinting. The structural stability of the vasculature construct was studied by observing the materials' erosion or degradation for 2 weeks of perfusion. Additionally, the cell adhesion and proliferation were quantitatively measured using CCK-8 kit method described in section 2.6.

2.9 Bioactive functionalization of the construct and in vitro hemodynamics.

The angiogenic peptide sequence with chemically functional group thiolated KLTWQELYQLKYKGINH₂ was designed and prepared to mimic the VEGF protein. Typically, a cysteine amino acid at the N- terminus of this sequence was introduced to allow its further reaction with the methacrylate group. Peptides were obtained with more than 95% purity according to the high-performance liquid chromatography (HPLC) profile provided by the manufacturer (GenScript). After mixing with the GelMA/C bioink (≈ 7.8 in PBS), VEGF peptide (50 ng/mL) was conjugated onto the material via Michael addition reaction.

To mimic the surrounding fluid present *in vivo*, a custom-designed flow bioreactor system was utilized for incubating 3D bioprinted vascular constructs in dynamic culture over the whole experiment period. The constructs (vasculature embedded in GelMA hydrogel) were placed in the PDMS chamber, and culture medium was perfused using a digital peristaltic pump (Masterflex, Cole-Parmer) at a certain flow rate. To choose an appropriate flow rate, the ranges from 1 mL/min to 100 mL/min were investigated. A fluid reservoir provided the culture medium for circulation, and a port for gas exchange, 5% CO₂/95% air was also included. For the fabrication of the PDMS chamber, PDMS (Sylgard 184, Dow Corning) base was mixed with a curing agent in a 10:1 weight ratio to make PDMS precursor. After removing air bubbles via centrifugation (5000 rpm, 10 min), the perfusable chamber was fabricated by curing at 80 °C for 1 h. The convective forces provided by creep flow, as the medium flows through the vasculature, facilitate an efficient transferring of nutrients and oxygen. Static culture served as the control and was identically operated in tissue culture

wells. For the burst pressure analysis, the printed vasculature was maintained in perfusion chambers with one end sealed, and the other end attached to a differential pressure gauge. The vasculature was filled passively with PBS until failure. To perform this dynamic development in the vasculature, representative mechanical parameters, specifically wall tensile stress (T) in the circumferential direction, were calculated using Laplace's Law [22].

$$T = \frac{Pr}{h}$$

(Equation. 2),

where h is the thickness of the vascular wall (mm), r is the vessel radius (mm), and P is the fluid pressure.

The average shear stress (τ) applied over the cell surface cultured in constructs was calculated using Poiseuille's law in case of laminar flow [22, 30].

$$\tau = \frac{4\mu Q}{\pi r^3}$$

(Equation. 3),

where τ is the average shear stress over the cell surface (dyn/cm^2), μ is the viscosity of the perfusing fluid (culture medium, $8 \times 10^{-4} \text{ Pa}\cdot\text{s}$ at 37°C), and Q is volumetric flow rate (mL/s).

2.10 Vascular permeability.

Diffusional permeability was quantified by perfusing culture media through the channel of 3D bioprinted vasculature [13, 31]. Texas red-labeled 70-kDa dextran ($25 \mu\text{g/mL}$, Thermo Fisher Scientific) was injected at a rate of $20 \mu\text{L/min}$ for 5 min, and then $1 \mu\text{L/min}$ for 2 h. The diffusion profile was detected and recorded every 10 min using a confocal microscope. The diffusional permeability of dextran was calculated by quantifying changes of fluorescence intensity over time using the following equation [13, 31]:

$$P_d = \frac{1}{l_1 - l_b} \left(\frac{l_2 - l_1}{t} \right) \frac{d}{4}$$

(Equation. 4),

P_d is the diffusional permeability coefficient, I_1 is the average intensity at an initial time point ($t=0$ min), I_2 is an average intensity after a different time interval, I_b is background intensity (before injecting dextran), and d is the channel diameter.

2.11 Vasoactivity and contractile matrix deposition.

To quantify the nitric oxide produced by 3D printed vasculature, after 1 day and 2 weeks of perfusable culture, the samples with 5 mL medium were maintained at -20 °C. After freezing and thawing for three cycles, proteins were removed using a microcentrifuge at 10,000 rpm for 5 minutes. Total nitrite in each sample was assessed using a Greiss reagent assay kit (Pierce). Samples were normalized to pure medium controls. To assess the endothelium-dependent vasodilation response, 1 μ M acetylcholine (Sigma) was added to the flow chamber for performing an acetylcholine response curve. To evaluate the effect of eNOS inhibition on the vasodilation, a final concentration of 3.2 μ M L- N^G-Nitroarginine methyl ester (L-NAME, Sigma) was added to the medium for 20 min. The construct was observed by optical microscopy (Mu800, Amscope) with a video camera, and the outer diameter change was analyzed using Image J. Changes in outer diameter are expressed as percent change from the baseline outer diameter. To assess the protein deposition of smooth muscle cells, the total collagen content was measured via Sirius red method. The suspension was dried and then incubated in Sirius red solution (0.1% Sirius red in picric acid) for 1 h, and then washed in 5% acetic acid. The resulting precipitate was dissolved in 0.1 M NaOH for 30 min. The OD was measured at 550 nm and the measurements were compared to collagen standards.

2.12 Fluorescent imaging and reverse transcription polymerase chain reaction (rt-PCR) analysis.

Fluorescent imaging and rt-PCR were used to investigate 3D vasculature development in vitro for each condition. Printed constructs were fixed with 10% formalin for 20 min, permeabilized in 0.1% Triton X-100 for 30 min and blocked in 2% BSA for 2 hours. To confirm the endothelium structure, double staining of F-actin and nuclei was performed using the method above. For immunostaining, the constructs were incubated with primary antibodies at 4 °C overnight. The following primary antibodies were used for staining: mouse monoclonal anti-platelet endothelial cell adhesion molecule-1 (PECAM-1/CD31) antibodies (Abcam) and rabbit monoclonal anti-alpha smooth muscle actin (α -SMA) antibodies (Abcam). After incubation with primary antibodies, chicken anti-mouse IgG-TR (Santa Cruz Biotechnology) and goat anti-rabbit IgG H&L Alexa Fluor® 488 (Abcam) as secondary antibodies were added and incubated for 1 h, respectively. Fluorescence images were observed using a confocal microscope.

The gene expressions of 3D bioprinted vasculature involving smooth muscle differentiation and vasculogenesis were quantitatively analyzed by rt-PCR, including vWf, CD31, myocardin, and α -SMA. After incubation for 1 and 2 weeks, the isolation of total RNA was performed on the samples using Trizol reagent. Total RNA purity and concentration were determined using a microplate reader (OD 260/280 nm within 1.8–2.0). RNA samples were reverse-transcribed to cDNA by using a Prime- ScriptTMRT reagent Kit (TaKaRa). RT-PCR was then performed on a CFX384 Real-Time System (BIORAD) by using SYBR Premix Ex

Taq™ (TaKaRa) according to the manufacturer's protocol. The gene expression level of target genes was normalized against the housekeeping gene GAPDH. The relative gene expression level was normalized against the control samples to obtain relative gene expression fold values and calculated via the 2-delta delta cycle-threshold (2- $\Delta\Delta$ Ct) method. Primer sequences are as follows: GAPDH, forward primer 5'-GGAGCGAGATCCCTCCAAA-3' and reverse primer 5'-GGCTCCCCCCTGCAA-3'; vWf, forward primer 5'-CACCATTCAGCTAAGAGGAGG-3' and reverse primer 5'-GCCCTGGCAGTAGTGGATA-3'; CD31, forward primer 5'-GAGTCCTGCTGACCCTTCTG-3' and reverse primer 5'-CACTCCTCCACCAACACCT-3'; myocardin, forward primer 5'-CTCGGCTTCCTTTGAACAAG-3' and reverse primer 5'-CTTCCAGAGAATCCATCCA-3'; α -SMA, forward primer 5'-CTGGGGTATTGGGGGCATC-3' and reverse primer 5'-CTGTTCCAGCCATCCTTCAT-3'. *In vivo* biocompatibility and biodegradation evaluation. 8-week-old, female, C57BL/6N mice were used to evaluate biocompatibility. All of the animal experiments were approved by the Institutional Animal Care and Use Committee of the National Heart, Lung, and Blood Institute. A method of random and blinding group allocation was applied to the animal experiments. Disc-shape GelMA/C hydrogels (300 μ L) were prepared, using similar printing conditions, and equilibrated overnight in PBS. Then the hydrogel constructs were surgically implanted into the subcutaneous space on the dorsal region of the mouse. At predesigned time points, post-implantation, the animals were euthanized and the implants, together with surrounding tissue, were harvested, fixed in formalin, and processed for histological analysis. Hematoxylin and eosin (H&E) staining was performed to evaluate the *in vivo* biocompatibility of the materials. Images were also taken to observe the degradation of the implanted hydrogels.

2.13 In vivo implantation of 3D bioprinted vasculature.

The *in vivo* development of the 3D bioprinted vasculature was evaluated using a xenograft model of transplantation into six-week-old female NOD-SCID IL-2 receptor gamma null (NSG) mice. All the animal experiments were approved by the Institutional Animal Care and Use Committee of the National Heart, Lung, and Blood Institute. A method of random and blinding group allocation was applied to the animal experiments. Each NSG mouse was subcutaneously implanted with the printed cell-laden vasculature, 10 mm diameter \times 3 mm thickness in size. All constructs were prepared using the protocols outlined above in a sterile condition and then surgically implanted into lateral incisions on the dorsal region. At weeks 1, 2, 4 and 6 of post-implantation, animals were euthanized and the specimens, along with the adjacent tissues, were collected for further examination.

2.14 Histology and immunohistochemistry.

Samples were fixed with 10% neutral buffered formalin and processed by freezing sectioning method. The sliced samples (10 μ m) were stained with H&E through visualizing cell nuclei and cytoplasm to observe cell distribution and development in the 3D printed blood vessels. Additionally, Masson's trichrome staining was used to identify muscle and collagen fibers. To characterize vascular maturation *in vivo*, sections were probed with specific antibodies for the immunofluorescence evaluation. Primary antibody against the

endothelial cell marker, monoclonal mouse anti-human CD31 (Dako, M0823) followed by secondary antibody labeled with Rhodamine (Jackson ImmunoResearch), were used to detect the endothelial cells and the lumen formation. For smooth muscle immunofluorescence, rabbit polyclonal anti-human α -SMA was used as the primary antibody, and FITC-conjugated secondary antibody (Jackson ImmunoResearch) was stained to visualize images. All the fluorescently stained sections were counterstained with Hoechst 33258. Fluorescence images were observed using a confocal microscope.

2.15 Statistical analysis.

All data are presented as the mean \pm sd (standard deviation). A one-way analysis of variance (ANOVA) with Student's t-test was used to verify statistically significant differences among groups, with $p < 0.05$ being statistically significant (*, $p < 0.05$; **, $p < 0.01$; ***, $p < 0.001$).

3. Results and discussion

3.1 Preparation of the tailored bioink and resultant elastomeric hydrogel

A tailored bioink catechol modified gelatin methacrylate (GelMA/C, Fig. 1A) was formulated that could be crosslinked by oxidant for hydrogel formation and functionalized with bioactive factors. The GelMA/C was synthesized through a two-step chemical reaction, in which gelatin was successively reacted with methacrylic anhydride and dopamine (Fig. S1, Supplementary Figures). Gelatin and its derivatives have been widely studied as promising biomaterials for vascular regeneration due to their native RGD (arginylglycylaspartic acid) adhesion motifs and endothelial-responsive degradation mechanism with matrix metalloprotease 2 or 9 [6, 13, 32]. Both play an important role in tissue remodeling associated with various physiological or pathological processes such as morphogenesis, angiogenesis, tissue repair, and metastasis, among others [33, 34]. Modification of biomaterials with a catechol moiety, known as mussel-inspired chemistry, has been widely used for surface functionalization to improve biocompatibility and enhance adhesive properties; it is also used for hydrogel formation via oxidation or enzyme induction [35–38]. In biology, dopamine, which is a typical catecholamine, plays several important roles in the nervous, immune, and digestive system as well as blood circulation [37]. In chemistry, pH-induced oxidation of the catechol group (>7.5) can result in a transition of catechol into o-quinone, which subsequently forms catechol–catechol adducts [37]. Previous studies have demonstrated that catechol modified polymers can form hydrogels in the presence of oxidants, such as tyrosinase, H_2O_2 , sodium periodate ($NaIO_4$), and O_2 [35, 39]. Results also showed a trace amount of $NaIO_4$ is a preferred crosslinking agent due to its rapid gelation and high biocompatibility both *in vitro* and *in vivo* [35, 40].

According to previous studies [41], to confirm the high efficiency of carbodiimide coupling, the excessive methacrylic anhydride was reacted to ensure no active amines remained in the first step. Then bioinks with various dopamine contents were synthesized via a carbodiimide coupling reaction in (-NH₂)/GelMA (-COOH) mole ratios of 10:1 and 5:1. The methacrylate groups would further serve to immobilize the bioactive factors. ¹H NMR spectra confirmed the successful conjugation of dopamine onto GelMA by the presence of catechol proton

peaks at around 7 ppm (Fig. S2). Fig. 1B shows UV-Vis spectra of GelMA/C and its oxidization product solutions (treated with NaIO_4). An absorption peak at 280 nm using dopamine standard solutions determined the content of catechol in the GelMA/C (Fig. 1C). After calculation, degrees of substitution (DS) of catechol was 6.0% for GelMA/C#1 and 13.3% for GelMA/C#2. GPC (water) results showed M_n (number-average molecular weight) of Gelatin was 4×10^4 with the polydispersity index (PDI) of 1.4, M_n of GelMA was 4.3×10^4 with PDI of 2.1, and M_n of GelMA/C was 4.6×10^4 with PDI of 2.3. In the current study, the gelation of the GelMA/C solution was induced by NaIO_4 with an equal molar ratio of the catechol group under neutral conditions ($\text{pH} \approx 7.8$) via generation of catechol–catechol adducts. Fig. 1D shows the photo images of the hydrogel formation (15 wt% GelMA/C#2 solution shown here). The color of the solution turned brown when adjusted to neutral due to the oxidation of oxygen in the air. After adding NaIO_4 solution, the self-standing hydrogel was formed immediately. Like any other hydrogels, the gelation time of GelMA/C depended on the DS of catechol, and solution concentration (Fig. S3). To obtain rapid gelation for 3D bioprinting, higher than 15 wt% solution was best for the GelMA/C with a different DS of catechol after numerous trials. Additionally, when the concentration of the bioink solution was higher than 30 wt%, it became too difficult to handle for the extrusion process, largely due to its high viscosity.

3.2 Physical characterization of GelMA/C hydrogel

To study the structural stability of GelMA/C hydrogel under cell culture conditions, the swelling properties of GelMA/C hydrogel (15 wt%) in Dulbecco's modified Eagle's medium (DMEM) at 37 °C were investigated by measuring weight changes during 7 days of incubation (Fig. 1E). The swelling of hydrogels reached equilibrium or saturation in 3 days. The GelMA/C#2 hydrogels with a high DS of catechol exhibited a lower swelling capacity with more rapid saturation compared to the low DS counterparts. This is due in large part to the higher degree of crosslinking. Any significant weight change or structural deformation was never observed over the entire experiment period, indicating the designed hydrogel would provide a stable environment for further cell loading.

Furthermore, the mechanical property of GelMA/C hydrogel was studied, which is a key functional parameter of tissue constructs for engineered blood vessels. In addition to structural support, withstanding contractile forces within engineered constructs is also very important for regulating hemodynamic behaviors and other physiological activities. The dynamic mechanical analysis was initially performed to test the rheological properties of GelMA/C hydrogels, which are affected by DS of catechol and concentration. The storage modulus measures the stored energy of dynamic mechanical analysis, representing the elastic mechanical strength of hydrogels. As a typical characterization of sol-gel transition, convinced data can be obtained in the linear viscoelastic region by varying loading conditions, such as frequency and strain. Fig. 1F depicts storage moduli (G') and loss moduli (G'') evolutions of hydrogels as a function of frequency. After stable gelation for 10 min, the G' for the GelMA/C hydrogel presented a substantial elastic response that was constantly greater than the G'' over the entire frequency range. As expected, a higher modulus of hydrogel would be obtained, and the modulus values changed from 10^2 Pa to 10^4 Pa when increasing concentration and DS of catechol. Additionally, all GelMA/C hydrogels

exhibited a stable elastic behavior without yield or failure when varying strain from 0.01% to 20% (Fig. 1G). By modulating graft polymer compositions or solution concentration, the elastic properties of the hydrogel could be adjusted to satisfy the requirement of cellular development and biomechanics, thus improving cellular functions and offering a better hemodynamic profile. According to previous experience, the hydrogel with a higher modulus has lower cell viability due to limited proliferation space and nutrient diffusion. The elastic modulus of the printed vessel was much lower than the actual value of the native vessel, thus it is unable to directly replace the native vessels. Considering that the engineering tissue scaffolds usually serve as a temporary supporting matrix, it is expected that the extracellular matrix will be deposited to form the native tissue environment of the blood vessel with the degradation of biomaterials after *in vivo* implantation.

In addition to serving as the crosslinking group, the catechol can be anchored to peptides and proteins due to its high binding affinity to diverse nucleophiles (e.g., amines, thiol, and imidazole), which may enhance cell or tissue adhesion [42]. The adhesion properties of the GelMA/C hydrogels were measured by recording detachment stress of hydrogel from the mouse muscle tissues, and photocrosslinkable GelMA hydrogels without catechol groups that have similar concentration and modulus served as the control. Fig. 1H revealed that the adhesion strength of GelMA/C hydrogels was significantly higher than that of GelMA hydrogel, and proportionally increased with increasing catechol contents. As the interaction between gel and muscle interface cannot be stronger than the inner force of hydrogel itself, the results suggested the catechol modified hydrogel possessed higher tissue adhesion compared to pure GelMA hydrogel. It is expected that the high adhesion ability would promote its integration with peripheral tissues.

3.3 3D bioprinting of vasculature and characterization

To create self-standing vasculature with a distinct smooth muscle layer and endothelial layer in an integrated manner, a custom-designed, 3D bioprinter with a coaxial needle system was developed to directly fabricate a tubular structure, and resultant stacked networks (Fig. 2A). Nozzle size controls the vessel's ID, and printing speed and extrusion flow rate affect the vessel's WT. It is known, WT that is too thin resulted in few encapsulated cells, while WT that is too thick might induce insufficient permeation of nutrients [43]. After optimization, a small-diameter vascular construct (~1 mm tube in diameter) was successfully produced using an 18G external needle and a 22G internal needle. The printing speed of the vasculature was maintained at 30 mm/s, and the extrusion flow rate of bioink for each syringe was set at 0.2 mL/s. The coaxial needle gauge can be easily replaced to achieve the fabrication of different-scale vasculature, which is similar to the reported study [43]. Although branching hierarchical trees are more biomimetic, a universal geometry-like model was chosen, which is widely used in many studies [13, 15, 44]. By replicating the essential information on structure, scale, orientation, and perfusion, it is suitable to explore a universal theory through hemodynamic experiments in the current study. Based on the material testing results above, 15 wt% GelMA/C#2 was selected for the bioprinting and testing of vascular constructs for the rest of this study due to its high mechanical strength and low viscosity among all samples. Generally, the bioprinting step was performed by extruding bioinks through the coaxial needle (Fig. 2B and Movie S1, Supplementary Movies). The bioink

mixture containing GelMA/C and human coronary artery smooth muscle cells (HCASMCs) flowed through the external needle, while the crosslinking slurry containing Pluronic F127 (30 wt%), NaIO₄, and human umbilical vein endothelial cells (HUVECs) flowed through the internal needle. Differing from the direct reaction between NaIO₄ and GelMA/C solution, the diffusion kinetics of NaIO₄ in the F127 slurry could affect the gelation process during 3D bioprinting. After numerous attempts, 5 mg/mL NaIO₄ was successfully used in the vascular bioprinting. When the oxidant encountered the bioink, the catechol groups of GelMA/C were rapidly crosslinked to form a tube with a core-shell structure. Fig. 2C shows a thick, vascular construct (20 layers and totally 2 cm in thickness) with regular lattice networks printed in a stackable manner, where HCASMCs were encapsulated into the shell layer, and HUVECs were encapsulated into the inner layer. As the construct was printed with a grid pattern, the supplement of nutrients and oxygen was determined by the thickness of the vessel wall. As such, it is expected that the cell viability would not be affected by the construct thickness.

The fugitive ink in the inner layer, Pluronic F127, at a certain concentration is a thermosensitive hydrogel material that is inert to cell adhesion. It can transform into a solution at low temperature or can be dissolved in water. With the fugitive slurry removal in medium, it is expected that HUVECs could migrate and adhere to the inner surface of the tube, although some of them were washed away with the fugitive removal. With 24 hours of incubation in medium, a hollow, self-standing “grid-like” vasculature construct with a bilayered cell structure was created after removing the fugitive slurry (Fig. 2D). Although the distance of each vessel was approximately 5 mm in the current images, the distance between the vessels could be optimized by changing the printing parameters, which depends on the real demand of muscularized vessels *in vivo*. In our preliminary trial without cells, the red dye was added to the fugitive ink to observe its removal process, thus confirming the 24 hours of the post-processing step. In other studies, a two-step method was applied to prepare the endothelium, where the F127 was rapidly removed by a cold solution flush, and then HUVECs were perfused into the lumen channel [13]. In the current design, a new method was used to ensure “one-step” endothelium formation. In the fluorescent images, a bifurcated blood vessel sectioned from the printed vasculature was observed, which is in the form of a bilayered lumen with smooth muscle layer (green, outer layer) and endothelial layer (red, inner layer) (Fig. 2E). The cross-sectional views of a representative segment of 3D bioprinted vasculature revealed vascular lumen formation. Similar to some previous studies [13, 15], the image quality for the printed vasculature with low magnification is limited to identify cell-cell junction, but the partially-enlarged scanning was conducted with high magnification in Section 3.7. It should be emphasized that, rather than traditional sacrificial template techniques, an integrative “fugitive-migration” manufacturing approach was presented where the fugitive slurry not only provided temporary support for the tubular structures during bioprinting but also assisted in crosslinking the GelMA/C bioink for hydrogel formation. More importantly, it may also anchor endothelial cells in the channel for endothelial cell layer formation and decrease the harmful effect of the oxidative crosslinker when compared to direct permeation of oxidant solution. A different technique was developed directly to form the endothelial layer, instead of indirectly with endothelial cell

perfusion, and the results were similar to, but more simple, than other endothelial cell perfusion studies [13].

After that, a cell-perfusion study was performed to observe dynamic flow behavior in the 3D printed vasculature, similar to blood circulation *in vivo* (Movie S2). Fig. 2F shows partial view images and their 3D optical maps (height images) of straight and bifurcated structures in the vasculature construct, which suggests the vasculature is perfusable and interconnected. Notably, after cross-printing, a geometric vessel network containing well-connected bifurcated structures at each joint interface was obtained, as opposed to a broken or leaky one, which is dependent on appropriate gelation time and effective gelation fashion to ensure the fusion of hydrogel. According to the data in Fig. S3, a 60 s gelatin time was allowed for achieving more stable intervessel junctions of the printed vasculature. Intervessel junctions that represent a step up in complexity are the basis for native vascular architectures which terminate in a single inlet and outlet. Fluid perfusion was performed through a red dye injection experiment to confirm the interconnected feature of the vasculature for mimicking blood circulation *in vivo* (Fig. 2G). Additionally, the cross-sectional morphological characteristics of the 3D bioprinted vascular constructs after lyophilization were observed using a scanning electron microscope (SEM) (Fig. 2H and I). The inner structure of a porous network in the vessel wall could be beneficial to the exchange of nutrients and gases in the cell-laden construct.

To maintain the long-term function of engineered tissue, the bioprinted construct must be able to maintain cellular homeostasis, self-renewal, and development. Both printing procedure (bioink, time, and printing modality) and post-printing culture may impact the cell viability. After bioprinting, the viability and proliferation of encapsulated HCASMCs and HUVECs were evaluated in the 3D printed vasculature. After 1, 3, and 7 days of culture, the Live-Dead staining (Fig. S4) indicated that the cell viability of both HCASMCs and HUVECs was preserved and not significantly influenced by the bioink materials (GelMA/C) and crosslinking agents (NaIO_4 and F127). Initial death of cells was observed on the first day due to the high shear force experienced during bioprinting and hydrogel encapsulation. However, it was significantly reversed with cell proliferation over the prolonged culture period. The results were normalized to the whole fluorescence area at day 1 (The initial cell viability was defined to 100%, so total cell viability was shown to be >100% after 7 days of culture). Moreover, cell metabolic activity (or proliferation) was investigated up to 7 days (Fig. S5). The results showed that both HCASMCs and HUVECs had increased metabolic activity in the 3D printed vasculature, although their activities are lower than the control groups (TCPs) due to the initial cell death and loss during vasculature fabrication. Additionally, it is likely that the reduced metabolic activities of these cells are a result of limited nutrient availability through the vascular constructs compared to the control conditions. The differences in metabolic activity between HUVECs and HCASMCs could be attributed to HUVECs being grown on the surface of the vasculature while HCASMCs were encapsulated into the vasculature hydrogel.

In previous studies, coaxial extrusion bioprinting succeeded in producing a vascular architecture with perfusable lumen [43, 45–47]. They all employed combination bioinks, composed of alginate and calcium chloride (CaCl_2), however, some reported weaknesses,

such as non-cell-adhesive properties, low proliferation abilities, and instability for long-term culture potentially limiting practical application [48, 49]. To demonstrate the advantages of the tailored bioinks, the cell adhesion at 6 h and proliferation at 7 days were investigated on different materials (Fig. 3A). These time points were selected based on the adhesion, spreading, and proliferation kinetics of endothelial cells in a previous study [50, 51]. Data showed HUVECs had a significantly better adhesion after 6 hours of culture, and a higher proliferation after 7 days of culture on the surface of GelMA/C hydrogels compared to alginate hydrogels. Moreover, the alginate hydrogels exhibited degraded structural stability after 2 weeks of perfusion (Fig. 3B) [52, 53]. Therefore, it is expected that the GelMA/C vessel would have high vessel durability, ensuring blood flow stability. These results suggested the new bioink possessed excellent characteristics, such as surface endothelization and mechanical stability, on fabricated vasculature constructs.

3.4 3D bioprinting of vasculature application: SLA-printed tissue matrix with coaxial extrusion-printed vasculature

Beyond the independent vasculature or blood vessels, the goal of bioprinting vasculature is to integrate it into engineered tissue matrices for maintaining metabolic functions. To verify this hypothesis, a dual 3D bioprinting platform was applied to generate a thick tissue matrix with embedded geometric vasculature (Fig. 3C). The uniqueness of the dual printing platform is to integrate two different bioprinting mechanisms (laser-based SLA and coaxial extrusion systems) into one [5]. As the extrusion-based bioprinting is limited by its low resolution, the high-resolution SLA printer was used to cure the gelatin-based bioink into the precisely patterned structure for tissue matrix fabrication using a similar method from our previous papers [54–56]. Human bone marrow-derived mesenchymal stem cell (hMSC)-laden GelMA hydrogel, fabricated by SLA bioprinting, was used as a universal tissue matrix model. During SLA bioprinting, the 3D bioprinted vasculature was embedded into the hMSC-laden constructs (Fig. 3D and E). Compared to the extrusion bioprinting, the advantage of SLA bioprinting, used here, is its ability to fabricate a 3D geometrical structure with a high resolution, although the negative effect of photocrosslinking on cell viability is a consideration. The embedded vasculature was designed to ensure uniform nutrient supplementation throughout the surrounding matrix.

In physiology, endothelium permeability in the vasculature is pivotal in the exchange of nutrients and gases with the peripheral tissue. Therefore, an effective permeability of printed vasculature is required to maintain the function of the vasculature itself and the peripheral engineered tissues. The Live-Dead cell staining was performed to investigate the hMSC viability in the hydrogel-based tissue matrices where the nutrient was provided by the adjacent vasculature (Fig. 3F). It is observed that cell viability significantly reduced beyond 300 μm from the vessel, suggesting a typical diffusive limitation. It indicates that the 3D bioprinted vasculature could provide efficient nutrient distribution through diffusing medium to hMSCs in the matrix for supporting peripheral tissue viability and development.

3.5 Bioactive functionalization and in vitro hemodynamic study

In addition to maintaining cellular bioactivity, it is crucial to provide an adequate biomicroenvironment including mechanical, chemical or biological signals for regulating

engineered tissue remodeling and maturation. The bio-functionalization, involving the incorporation of bioactive factors, enzymatic recognition sites, and adhesion factors among others, is a promising approach for directly activating cellular signal pathways during tissue development. In this study, the methacrylate groups in the tailored bioink can be chemically immobilized with bioactive factors as in a previously reported study [57]. Thiol-functional oligopeptides with sequences of KLTWQELYQLKYKGINH₂ derived from the knuckle epitopes of vascular endothelial growth factor (VEGF) were conjugated to the GelMA/C polymer (Fig. 4A and Fig. S6), which was verified as a highly effective factor for vasculogenesis [6, 58]. Compared to directly incorporating VEGF growth factor, the peptide not only delivers specific bioactive factors in a uniform manner for guiding vascular formation, but also addresses issues of prolonged tissue retention and sustainable bioactivity compared to traditional growth factor encapsulation. As a small number of peptides were added, obvious spectra signals by IR and NMR were not successfully detected, where only the strong signals of the gelatin were observed [6].

Blood vessels are normally exposed to two types of mechanical forces *in vivo*: (a) circumferential stress acting tangentially on the vascular wall and directly related to pressure and dimensions (diameter and thickness) of the vessel, and (b) shear stress acting in the longitudinal direction at the blood-endothelium interface and directly related to the flow-velocity profile. To investigate the functional characteristics of the perfusable vasculature in a fluid dynamic environment, the vascular constructs were installed in a PDMS chamber with a single inlet and outlet, connected to a customized bioreactor. The embedded vascular network in the GelMA hydrogel constructs was designed as an asymmetrically branched structure to ensure uniform perfusion throughout the tissue for transporting nutrients, oxygen, and waste materials (Fig. 4B and Movie S3). To minimize the mechanical effect of the mounted hydrogel on the vascular construct, a distinctly softer hydrogel (only 2×10^2 Pa) was used. Fig. 4C illustrates the circumferential stress and shear stress during the perfusion culture. The flow rate with respect to the fluid pressure was varied to observe the distensibility of the vasculature construct. Burst pressure was measured by pressurizing vasculature constructs with PBS until failure (Fig. 4D). Compared to static culture, dynamic culture increased the mechanical properties of the constructs after 7 days. Although the burst pressure result is slightly lower than the native vessel, the engineered vasculature is suitable for supplying the blood within the engineered tissue constructs when compared to other studies [8]. Considering the elasticity and stability of 3D bioprinted thin vascular walls, a flow rate of 6.0 mL/min was selected after initial screening, for the dynamic culture studies. Through calculation, the average tensile stress (T) of the vasculature was 16.7 kPa (~62.5 mmHg), and the average shear stress (τ) was approximately 8.2 dyn/cm², which is within the range (from 5 to 12 dyn/cm²) of shear stress in the microcirculation [26]. The Laplace and Poiseuille's law were applied here as typical equations to obtain the value of hydrodynamics according to previous studies [22, 30]. The vasculature constructs were perfused for 2 weeks to evaluate vessel maturation and vasoactivity over time.

3.6 Vascular permeability, vasculoactivity, and extracellular matrix deposition

Maintenance of the size-selective sieving property of endothelium is crucial for several physiological functions, including normal tissue-fluid homeostasis, angiogenesis, vessel

tone, and host defense, which involves two endothelial transport mechanisms, transcellular and paracellular pathways [59]. To further assess the permeability function of the printed vasculature, dextran was used as a macromolecular model to test the diffusional permeability (Fig. 4E). After the endothelial layer formation in the vascular channel, the permeability of medium diffusing from the vasculature to the thick tissue matrix was observed at approximately 4.67×10^{-5} cm/s; this is consistent with convective diffusion transport and metabolic conversion within the tissue. Additionally, compared to the bare channel (without endothelial layer), a significant (almost 4.5-fold) reduction in permeability and no significant leakage of the culture medium during perfusion into the vasculature channel was observed. It can be concluded that the endothelial layer is dominating transport resistance, like the way it governs the permeability of blood vessels *in vivo*.

Physiologically relevant reactions to vasoactive stimuli were studied by nitric oxide (NO) release and endothelium-independent vasodilation for 2 weeks of culture. The endothelium synthesizes and releases NO to maintain homeostatic function; endothelium-derived NO maintains a non-thrombogenic surface, prohibits leukocyte attachment, and promotes vascular relaxation under basal conditions [60, 61]. NO produced by endothelial cells was evaluated by assessing total nitrite concentration, the stable NO metabolism byproduct under physiological conditions (Fig. 4F). NO production of the 3D printed vasculature constructs after 3 days of culture was close to the value after 2 weeks, in static culture. The dynamic culture significantly improved the NO production of the constructs when compared to the static culture. To further explore the role of NO in the endothelium-dependent vasodilation in the vasculature constructs, the effect of vessel diameter on the responses to acetylcholine (an endothelium-dependent vasodilator) with or without L-N^G-Nitroarginine methyl ester (L-NAME, an inhibitor of endothelial nitric oxide synthesis (eNOS)), were investigated [62, 63]. After 1 μ M acetylcholine was added into the culture medium, the diameter of the vasculature constructs dilated about 5%, and maintained relative stability for vasodilation response over 2 weeks, suggesting sustained endothelial health of the 3D printed vasculature (Fig. 4G). The dose- response of acetylcholine from 0.01 to 100 μ M showed increasing vasodilation for 2 weeks; however, after pre-treating with L-NAME for 10 min, the vasculature constructs exhibited vasoconstriction in the same conditions (Fig. 4H). The results demonstrated that the vasodilation in response to acetylcholine was attributed to the NO release of the 3D printed vasculature. Additionally, collagen is the major matrix constituent of the vessel wall and is synthesized by smooth muscle cells [64, 65]. Therefore, total collagen synthesis was used to quantify the ECM protein deposition. Fig. 4I shows that collagen production was increased over 2 weeks of culture, and the collagen content under dynamic culture was significantly higher compared to static culture. It indicated that hydrodynamic culture or biomechanical stress could improve the ECM deposition of 3D printed vasculature constructs. In the previous study, Tranquillo et al. have also verified that mechanical forces play an important role in shaping the organization of the ECM when they applied a cyclic stretch or pulsed flow stretch to the grafts in a bioreactor [66, 67].

3.7 Vasculogenesis and contractile protein expression

For the functional development of 3D bioprinted vascular constructs, smooth muscle differentiation medium was added to induce HCASMC differentiation according to

ThermoFisher's protocol (HCASMCs will undergo a cessation of growth, a change in cellular morphology from slender stellate cells to enlarged rectangular or triangular shaped cells, and will express increased amounts of smooth muscle α -actin over the course of 2 to 8 days). After weeks of active perfusion, the structure of 3D bioprinted vasculature maintained good integrity under a shear stress of 8.2 dyn/cm². As shown in Fig. S7, these endothelial cells maintained their endothelial phenotype and distributed uniformly around the lumen characterized by expression of platelet endothelial cell adhesion molecules (PECAM-1/cluster of differentiation 31, CD31; makes up a large portion of endothelial cell intercellular junctions involved in leukocyte transmigration, angiogenesis, and integrin activation). Moreover, smooth muscle cells encapsulated in the outer layer exhibited the α -actin phenotype of smooth muscle (α -SMA, are highly conserved proteins that are involved in cell motility, structure and integrity, and a major constituent of the contractile apparatus). However, as our vascular construct diameter (> 1 mm) is larger than the resolution of confocal microscopy, it is hard to identify the confluent feature in the tile scanning model [13]. Thus, partially enlarged scanning with a high magnification was performed. In response to an integrated stimulus with angiogenic VEGF peptide and dynamic culture, Vinculin staining showed HUVEC growth was aligned with flow direction, which can be attributed to the shear stress (Fig. 5A and Fig. S8). Vinculin is a protein that couples, transmits, transduces, and regulates mechanical force between the cytoskeleton and adhesion receptors. F-actin and VE-Cadherin staining of HUVECs also revealed a confluent endothelium with intercellular junctions on the lumen surface of 3D bioprinted vasculature at day 7, and the HUVEC behaviors were similar to a previous study [13] (Fig. 5B and Fig. S8). Additionally, smooth muscle cells exhibited a switch to contractile phenotype (fusiform or elongated shape) with a strong axial alignment compared to the control, identified through co-staining of F-actin and α -SMA (Fig. 5C and Fig. S8). As mentioned above, the results further demonstrated the ECM alignment or contractile protein organization was determined by the mechanical stretch direction as evidenced previously [67].

To determine cellular functions in the 3D bioprinted vasculature, relative gene expression including endothelial function and smooth muscle differentiation was quantitatively measured by RT-PCR using GAPDH as a housekeeping gene. To deconvolute the mixing effects for co-culture, a pure cell population experiment was conducted. As shown in Fig. 5D and E, after 2 weeks of culture the von Willebrand factor (vWf, a blood glycoprotein) and CD31 expression in the VEGF peptide bonding groups showed increased vasculogenesis both in static and dynamic culture conditions. This demonstrates the ability of immobilized VEGF peptides to enhance signaling and bioactivity of endothelial cells and consequent vasculogenesis *in vitro*. Moreover, the vWf expression of the VEGF peptide bonding group in the dynamic culture is up-regulated 7-fold compared to the static culture. The expression of α -SMA and myocardin (Myo, a smooth muscle-specific transcriptional coactivator of serum response factor) has a similar tendency with the gene expression level of the dynamic culture group being much higher than the static culture counterpart (Fig. 5F and G). The results suggest that perfusion culture produces a beneficial effect for vascular smooth muscle formation by undergoing contractile force associated with hemodynamic pressure [20, 22, 68]. Additionally, it was also observed that the VEGF peptides promoted smooth muscle differentiation of HCASMCs to some extent. Overall, the VEGF peptides improved

vasculogenesis and smooth muscle differentiation of 3D bioprinted vasculature by providing a positive biological stimulus, while perfusion culture facilitated an efficient bio-environment, through both homogenous mass transport and biomechanical stimulation, improving matrix reorganization and vascular maturation.

3.8 In vivo GelMA/C hydrogel degradation and biocompatibility

Apart from ensuring cell viability and development throughout, *in vivo* biocompatibility of bioink, and their degradation byproducts, is necessary for tissue development and remodeling over long-term implantation. Additionally, another major consideration is the balance between implant degradation rate and neo-tissue formation. Subcutaneous implantation was performed on C57BL/6N mice to evaluate the biocompatibility of GelMA/C hydrogels. It is observed that the hydrogel disks slowly were degraded *in vivo* during the 16-week evaluation period, although complete disappearance did not occur during the experimental period (Fig. S9A–D). Compared to photo-crosslinked GelMA hydrogels (the hydrolysis and enzymatic cleavage of the amide bonds in the gelatin backbone), the GelMA/C showed a slower degradation due to the strong crosslinking of dopamine chemistry (Fig. S9E). The slow degradation is thought to be beneficial for structural stability of the 3D bioprinted vasculature, minimizing the risk of the walls thinning or weakening over the long-term process of vascular regeneration.

Furthermore, at different time intervals, the host tissues surrounding the hydrogel disks were collected and then stained by H&E to examine the inflammatory response. H&E is typically a pathological evaluation for the inflammatory response of implants. After the first week, only the outer margins of the hydrogel were infiltrated by immune cells, and few monocytes, indicating an acute inflammatory reaction in the initial stage (Fig. S9A). By 4 weeks, the number of infiltrating cells mildly increased, including monocytes and macrophages, and a fibrosis layer appeared around the hydrogel (Fig. S9B). By 8 weeks, the number of infiltrating inflammatory cells was dramatically reduced, as had the extent to which the material had been infiltrated through the center of the hydrogel. A fibrosis capsule was formed, and completely encircled the hydrogel (Fig. S9C). At 16 weeks post-implantation, the immune cells had almost disappeared, and only fibrosis was observed, indicating the appearance of an appreciable inflammatory response (Fig. S9D). Additionally, there was none of any obvious tissue necrosis, edema, hyperemia, hemorrhaging, and muscle damage observed during the experimental period. Histological analysis showed the materials could be degraded slowly, and the tissue reaction *in vivo* was consistent with a typical foreign body reaction with no severe chronic inflammation, implying that the synthetic material may be suitable for *in vivo* applications.

3.9 In vivo vasculature development in the immunodeficient murine model

To determine whether the constructs could support implanted cell survival and functionalization, the cell-laden vascular constructs were immediately implanted into NOD-SCID IL-2 receptor gamma null (NSG) mice after bioprinting. The subcutaneous tissue or hypodermis, consisting primarily of loose connective tissue and lobules of fat, contains a large number of blood vessels, suitable for the implantation of soft vascular constructs. To observe the “fugitive-migration” behavior and implanted cell development in situ, there was

neither performed the *in vitro* culture before implantation nor directly sutured the constructs with the host's circulation. In addition to developing engineered blood vessels, the *in situ* vascular development is significantly important for complex tissue regeneration.

Seeing that the hollow vasculature was easily crushed, after bioprinting the vasculature was mounted with a supporting GelMA hydrogel on the opposite side of muscle tissue), and then implanted into the dorsal region of the mice. The mount may help stabilize the vasculature *in vivo*, avoiding distortion from vigorous activity. It is noted that to eliminate the non-specific reaction of cell staining, human-specific vascular markers were used, and the images were taken from the implant parts without any host tissue in the following slide examination. Fig. 6A shows the cross-sectional schematics of a single-vessel cut from the bioprinted vascular construct, including HCASMC-laden GelMA/C hydrogel (shell) and HUVEC-laden fugitive slurry (core). At 1-week post-implantation, histological examination by H&E staining revealed that the vascular constructs were stratified *in vivo* and the fugitive slurry was embedded in the lumen of the vasculature (Fig. 6B). The HCASMCs and HUVECs were extruded from the shell and core layers of the axial bioprinter, respectively; therefore, HCASMCs (blue arrow) and HUVECs (red arrow) appear in the different regions of the images. Since the thin vascular wall slipped or was easily broken during the staining process, it was unable to observe the intact vascular structure in the images. Fig. 6C shows H&E, Masson, and immunofluorescence (Hoechst, blue; CD31, red; α -SMA, green) stained images of the implanted vascular constructs with 10x cross-sectional view at week 1, 2, 4, and 6. All of these images show the same area that the cuts were performed in Fig 6 A and B. By the second week, a bilayered vascular structure surrounded by host murine fibrosis tissue was observed, accompanied by the gradual removal of the fugitive slurry in the lumen of the vasculature. Small vessels (red circles) containing erythrocytes emerged, and connected the host cell layer with the implanted construct. These infiltrated murine blood vessels indicated functional connections, or connections occurring between 3D bioprinted vasculature and host murine vasculature. Although there is no inflammation response for the implants in the NSG mice, the fibrosis based on the native healing mechanism could involve stimulated fibroblasts laying down connective tissue. Thus, proliferation and activation of fibroblasts deposited extracellular matrix onto the surface of 3D bioprinted vasculature for the formation of surrounding connective tissue (tunica adventitia). The artificial structure (dual layer) of the 3D bioprinted vasculature was developed to replicate a mature and biomimetic vessel with three native layers (although the tunica adventitia was formed by the migrated fibroblasts of mice). After 4 weeks, numerous small vessels (red circles) were uniformly distributed throughout the host fibrosis tissue, and implanted vascular constructs highlighted using H&E staining. Masson staining also verified host murine fibrosis (or connective tissue, blue fibers) surrounded the 3D bioprinted vasculature (muscle fibers, red fibers and pick cytoplasm). Compared to the results at 2 weeks, a compact murine fibrosis tissue occurred, and further fused, with the implanted smooth muscle layer. After 6 weeks of implantation, a typical remodeling process of the vascular structure containing distinct connective tissue layer (tunica adventitia), smooth muscle layer (tunica media), and endothelium (tunica intima) was observed. Moreover, the inset images also verified that some small capillaries existed in the 3D bioprinted vasculature, which sprouted from the host vessel of the mice. The host vessel invasion provided a beneficial microenvironment for

implanted vasculature through an exchange of nutrients and oxygen. It is deduced that the host murine fibroblasts were involved in matrix reorganization, and enhanced muscularized vessel maturation in the vascular development illustrated by the Masson staining results.

Overall, these *in vivo* results uniquely demonstrated the *in vivo* development process of 3D bioprinted vasculature, including the formation of functional vessels and *in vivo* autonomous connection [33, 34]. As mentioned above, the current vascular construct is not robust enough to directly connect the blood circulation, although the *in vivo* perfusion is a perfect model for the current system. The 3D bioprinted vasculature will be further optimized in the future study to satisfy these requirements in potential clinical application. It is expected that this study will advance the success of vascular bioprinting that enables growth and maturation of thick tissues with critical defects through the support of vascularization driven by natural processes in the body.

4. Conclusions

In summary, a novel bioink and manufacturing tactic associated with a comprehensive post-bioprinting process were developed to create vascular constructs with necessary architectures (straight and bifurcated tubular structure). The major advantage of this method over others is the 3D bioprinted vasculature replicates biomimetic vessel structures, essentially consisting of smooth muscle and endothelium, with the goal of addressing crucial challenges of a small diameter blood vessel network. Biofunctionalization and dynamic stimuli dramatically improved vascular remodeling of both smooth muscle and endothelium. Favorable biocompatibility *in vivo* ensured the implantation safety of the material for further clinical application. Studies of 3D bioprinted vasculature development in immunodeficient mice verified the effectiveness of the design. Although more optimization is required to improve the bioprinted vascular construct, advanced achievements in creating biomimetic, functional vasculature show significant potential toward eventually fabricating a complicated tissue/organ for clinical transplantation.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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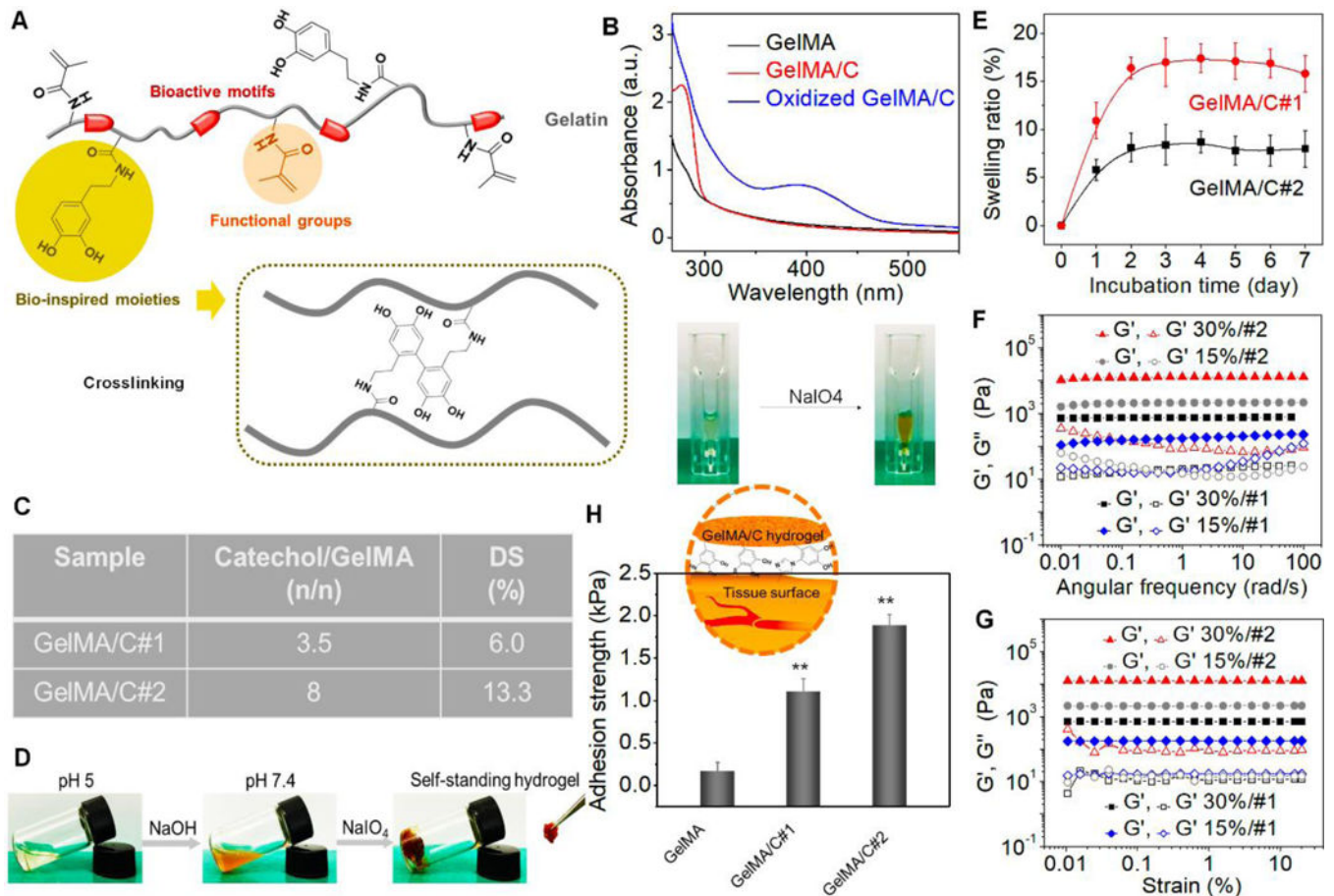


Fig. 1. Characterization of GelMA/C bioink and resultant hydrogel.

(A), Schematics of a GelMA/C copolymer structure and its oxidative crosslinking process, highlighting the key functional groups. (B), UV-Vis spectra of the GelMA/C solution (red line) that underwent oxidation (blue line), a GelMA solution serves as the control (black line). The colorless solution became brown after oxidation. (C), Degrees of substitution (DS) of catechol (GelMA/C#1 and #2) determined by the standard curve method. (D), Photo images of the GelMA/C gelation process. The GelMA/C solution (colorless) at pH 5 was adjusted to pH 7.8 (brown, sol), and further crosslinked to form the self-standing hydrogel (brown, gel). (E), Structural stability of the GelMA/C hydrogel in an aqueous environment through a swelling test after 7 days of incubation. (F), Dynamic mechanical analysis of the GelMA/C hydrogels. Storage moduli (G') and loss moduli (G'') evolution of the hydrogels with different DS and concentration (different colors) was measured in a frequency sweep mode. Filled symbols are storage moduli G' , and open symbols are loss moduli G'' . (G), Modulus-strain plots of the GelMA/C hydrogels. G' and G'' evolutions of the hydrogels with different DS and concentration was measured. (H), Adhesion properties of the GelMA/C hydrogels tested by detachment stress of hydrogel from the mouse muscle tissues. Compared with the GelMA hydrogel without catechol group, the GelMA/C hydrogels showed a significant increase in tissue adhesion ability due to a high binding affinity to peptides and proteins of the tissues; the mean \pm sd., $n = 9$, $**P < 0.01$.

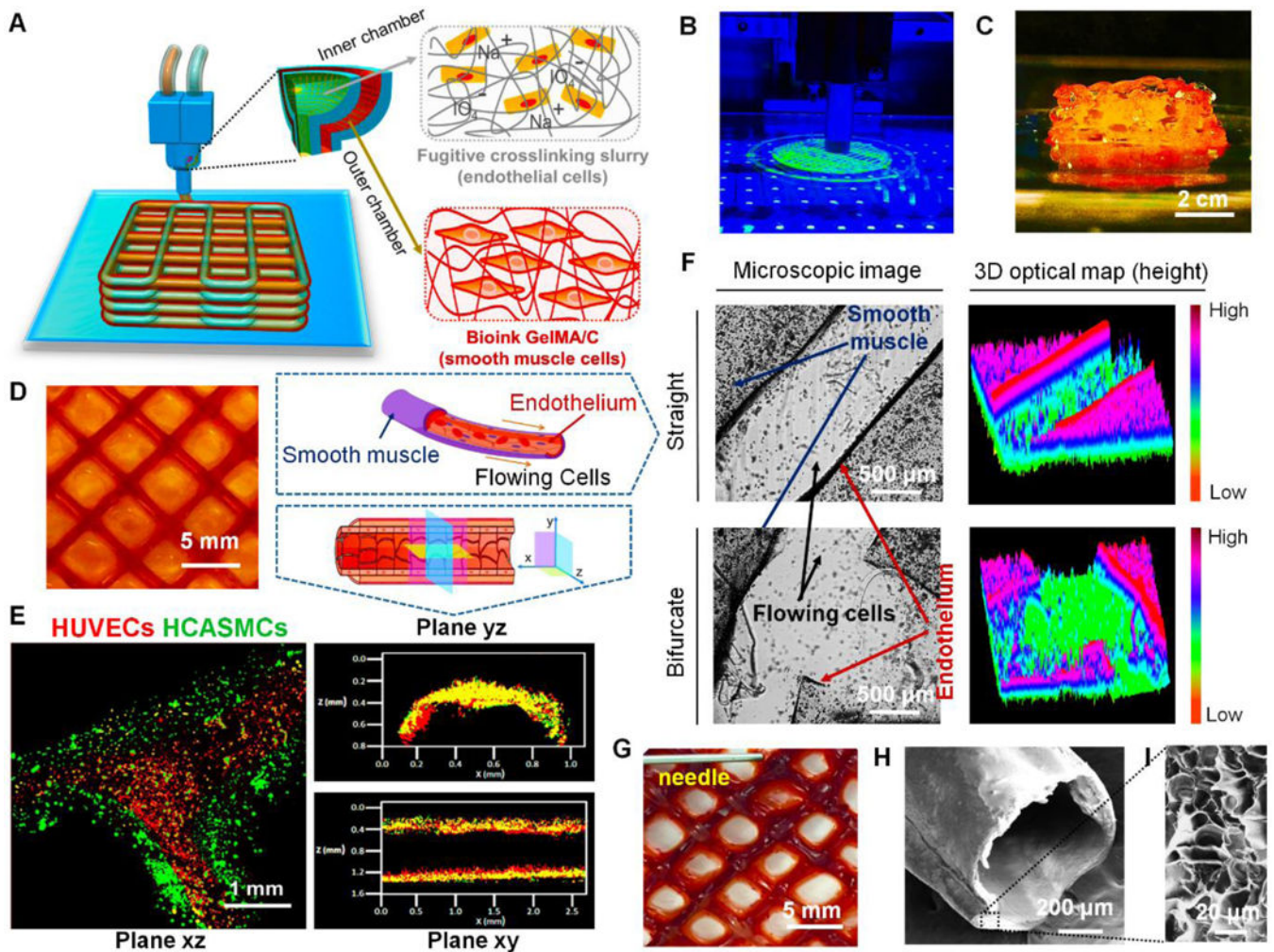


Fig. 2. 3D bioprinting of vasculature construct.

(A), an illustration of basic patterning of 3D vascular architecture fabricated by a coaxial extrusion bioprinter system. The “core-shell” filament consists of two parts: (i) the inner chamber contains fugitive crosslinking slurry and endothelial cells, (ii) the outer chamber contains GelMA/C bioink and smooth muscle cells. (B), a photo image of 3D bioprinting of vasculature on the self-made bioprinter. (C), the side-view image of the 3D bioprinted self-standing vascular construct with 2 cm height. (D), the top-view image of 3D bioprinted vasculature with hollow and interconnected structure (2 layers) after 24 hours of perfusion culture. (E), structural illustration and fluorescence images of 3D bioprinted cell-laden vasculature in three different planes. Endothelial cells were colored red, while smooth muscle cells were colored green. (F), microscopic images and their 3D optical maps (height) of 3D bioprinted vasculature (blood vessel) in straight and bifurcated regions, including smooth muscle layer and endothelium. Flowing cells were observed in the hollow channel after cell perfusion. (G), a photo image of blood plasma perfusion mimicked through a dye injection experiment. (H), cross-sectional morphology of 3D bioprinted vasculature taken by SEM. (I), enlarged morphology of the cross-sectional surface of 3D bioprinted vasculature taken by SEM.

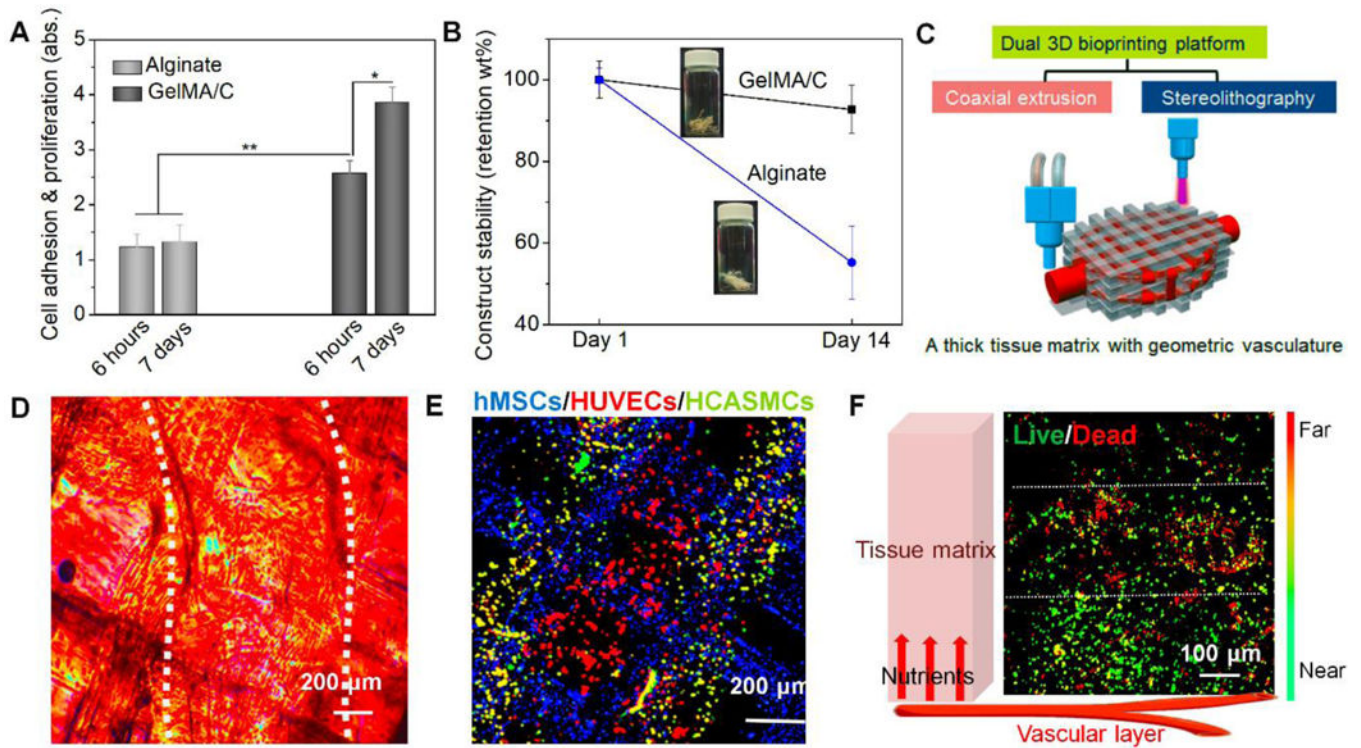


Fig. 3. 3D bioprinting of alginate-based vasculature and 3D bioprinting of tissue matrix with the vasculature.

(A), cell adhesion and proliferation of vascular constructs fabricated by GelMA/C and traditional alginate; the mean \pm sd., n = 9, *P < 0.05, **P < 0.01. (B), structural stability of vascular constructs fabricated by GelMA/C and traditional alginate. (C), schematics of a dual 3D bioprinting platform (coaxial extrusion and stereolithography) to be applied to generate thick, vasculature embedded tissue matrix, where the vascularized tissue is fabricated with vasculature in a “sandwich” manner. (D), microscopic images and (E), a fluorescence image of a 3D bioprinted cellularized tissue matrix. The white dotted lines in D indicate the position of 3D printed vasculature. Dual-layered vasculature (smooth muscle, green; endothelium, red) was embedded into an MSC-laden tissue matrix (grid, blue). (F), nutrient supplementation assay of encapsulated MSCs in the vascularized tissue using Live-Dead staining. Near/Far dedicates the distance away from the vasculature in the schematic. White dotted lines separate into three zones, i.e., death zone without enough nutrients, transitional zone, and survival zone with enough nutrients. It showed an effective supplementation of nutrients for cell viability related to diffusion.

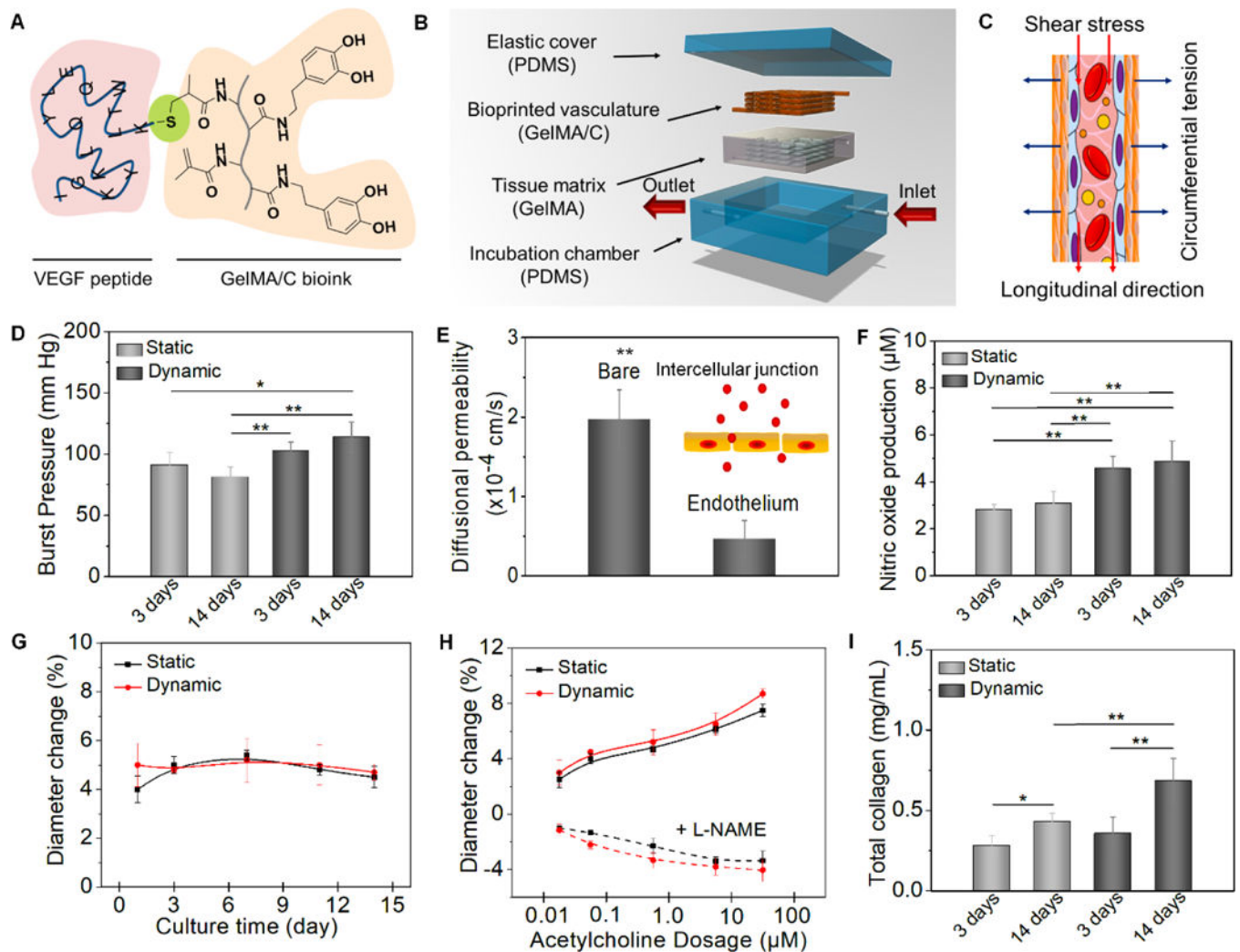


Fig. 4. Hydrodynamics, vasculoactivity, and extracellular matrix deposition.

(A), schematics of chemical immobilization of VEGF peptides onto GelMA/C bioink. A Michael addition reaction between methacrylamide and thiol-functional oligopeptide was performed for ink biofunctionalization in current design. (B), an illustration of dynamic perfusion culture. The vasculature with one inlet and one outlet was embedded into a tissue matrix (GelMA hydrogel as a supporting material) and then placed into a customized PDMA chamber. (C), an illustration of the circumferential stress and shear stress during the perfusion culture. (D), the burst pressure of 3D printed vasculature constructs under static and dynamic culture; the mean \pm sd., $n = 9$, * $P < 0.05$, ** $P < 0.01$. (E), quantification of barrier properties confirmed by the comparison of a bare channel (without endothelial lumen) and endothelial lumen. Both contain smooth muscle cells. By mimicking native endothelial functions, vascular permeability originates from the intercellular junction; the mean \pm sd., $n = 9$, ** $P < 0.01$. (F), nitric oxide production of 3D printed vasculature constructs under static and dynamic culture; the mean \pm sd., $n = 9$, ** $P < 0.01$. (G), vasodilation response to $1 \mu\text{M}$ acetylcholine of 3D printed vasculature constructs under static and dynamic culture over 2 weeks; the mean \pm sd., $n = 6$. (H), dose-dependent vasodilation response to acetylcholine of

3D printed vasculature constructs and their constriction induced by pre-treat of L-NAME under static and dynamic culture; the mean \pm sd., n 6. (I), total collagen synthesis of 3D printed vasculature constructs under static and dynamic culture; the mean \pm sd., n 9, *P <0.05, **P <0.01.

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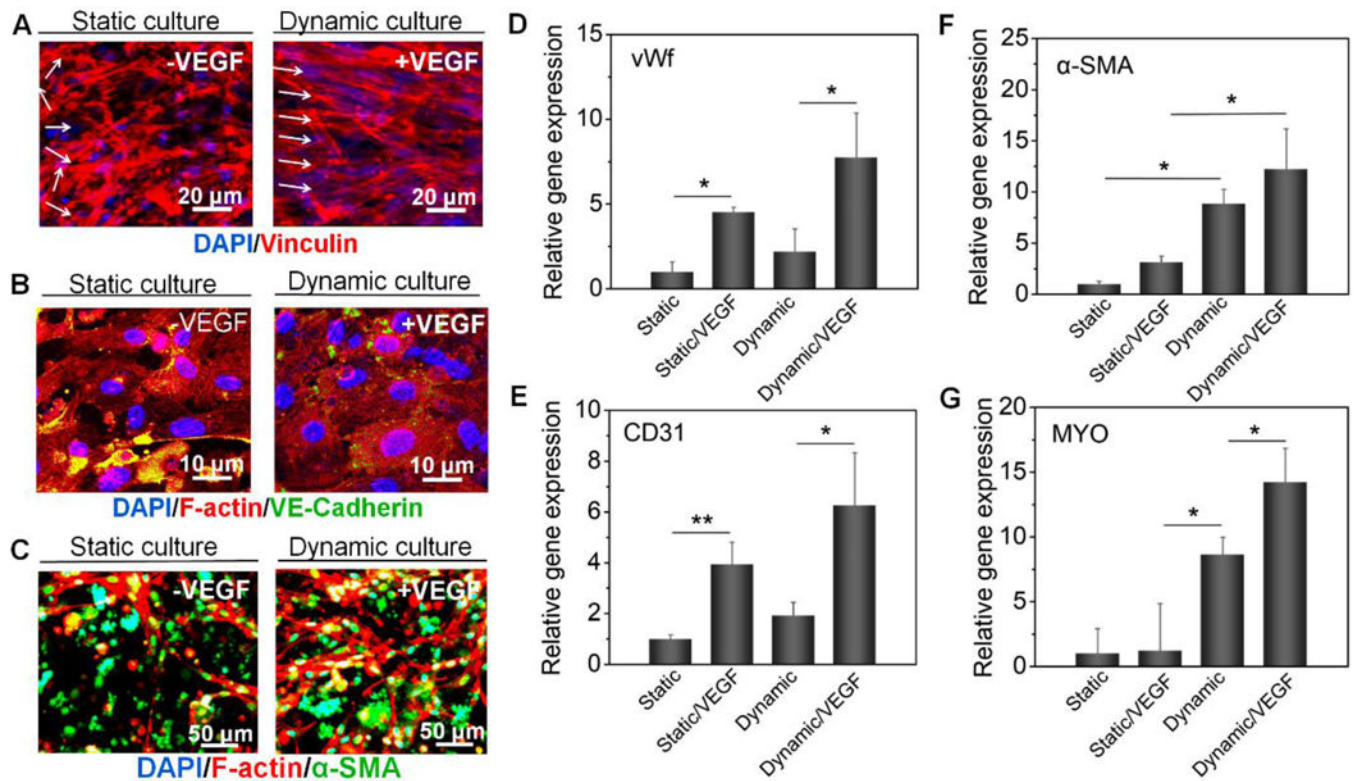


Fig. 5. Vasculature development study *in vitro*.

(A), immunofluorescence images of the endothelium (spreading and orientation) of 3D bioprinted vasculatures (vinculin, red; DAPI, blue) including +VEGF/dynamic group and -VEGF/static group. Images of other groups are shown in the supplementary file. (B), immunofluorescence images of the endothelium (intercellular junctions) of 3D bioprinted vasculatures (VE-cadherin, green; F-actin, red; DAPI, blue) including +VEGF/dynamic group and -VEGF/static group. Images of other groups are shown in the supplementary file. (C), immunofluorescence images of the smooth muscle of 3D bioprinted vasculatures (α -SMA, green; F-actin, red; DAPI, blue) including +VEGF/dynamic group and -VEGF/static group. Images of other groups are shown in the supplementary file. (D-G), Relative gene expression of vWf (D), CD31 (E), α -SMA (F), and MYO (G) with univariate analysis, including +VEGF/dynamic group, -VEGF/dynamic group, +VEGF/static group, and -VEGF/static group; the mean \pm sd., n = 9, *P < 0.05, **P < 0.01. The experiments were repeated 4 times. Results suggested VEGF peptide improved vasculogenesis and smooth muscle differentiation of the 3D bioprinted vasculature, while perfusion culture facilitated an efficient bio-environment to improve vascular maturation.

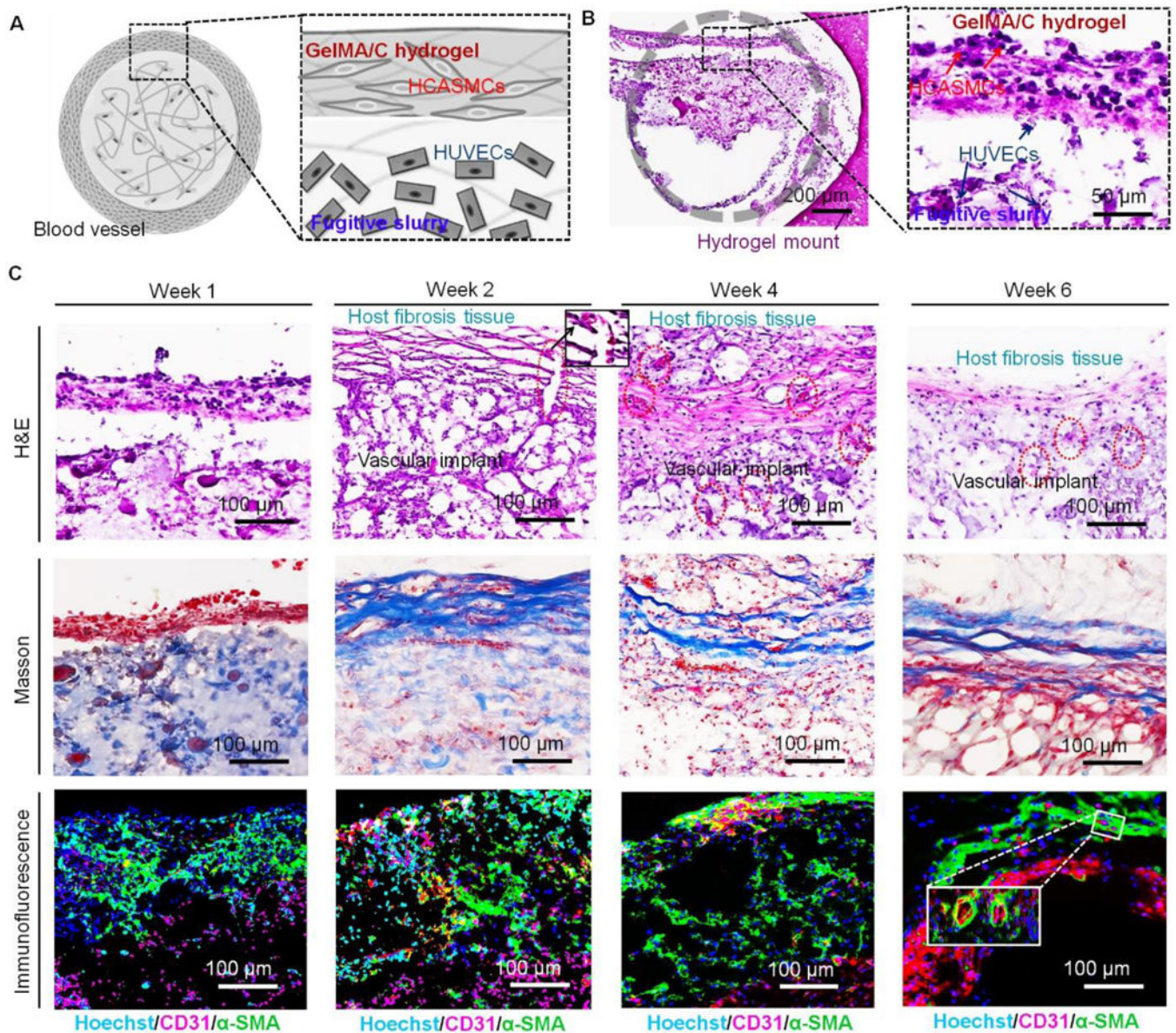


Fig. 6. *In vivo* development of 3D bioprinted vasculature in NSG immunodeficient mice. (A), cross-sectional schematics of a single-vessel that was cut from the bioprinted vasculature construct, including HCASMC-laden GelMA/C hydrogel (shell) and HUVEC-laden fugitive slurry (core). (B), H&E stained sections of the cell-laden vasculature with 1x and 20x cross-sectional view harvested at week 1. Gray circle indicates a round vessel lumen encircled by GelMA hydrogel mount. A distinct interface was observed between HCASMC-laden GelMA/C hydrogel and HUVEC-laden fugitive slurry. Some HUVECs adhered to the surface of GelMA/C hydrogel. (C), H&E, Masson, and immunofluorescence (Hoechst, blue; CD31, red; α -SMA, green) stained images of the implanted vasculature construct with 10x cross-sectional view at week 1, 2, 4, and 6. After 2 weeks of implantation, the vasculature was surrounded by the activated host fibroblasts. Vessels (red circles) containing erythrocytes connected the host cell layer with the implants highlighted with an enlarged image (black square). Masson stained images show the host murine fibrosis (or connective

tissue, blue fibers) surrounding the 3D bioprinted vasculature. Immunofluorescence images show the structural feature of the vasculature constructs, which is endothelium (red) surrounded by smooth muscle (green). After 6 weeks of implantation, a hollow and mature vessel structure with compact smooth muscle and endothelium was gradually generated. The inset image shows some small vessels generated in the bioprinted vasculature construct.