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Vascular microphysiological systems (MPS): biologically relevant and potent models

Vascular microphysiological systems (MPSs) are gaining increasing significance due to advancements in microvascular patterning, three-dimensional organization, and both cellular and acellular modelling. These improvements have enhanced their biological fidelity, making vascular MPS powerful tools for investigating vascular biology in relation to key physical parameters such as flow, stretch, and permeability. In parallel, their translational relevance continues to grow, with promising applications in disease and cancer modelling, immunological studies, and preclinical drug screening. Together, these features position vascular MPS as versatile platforms bridging basic science and biomedical innovation.

As featured in:



See Ryuji Yokokawa *et al.*,
Lab Chip, 2025, **25**, 4221.



Cite this: *Lab Chip*, 2025, **25**, 4221

Vascular microphysiological systems (MPS): biologically relevant and potent models

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Extensive research has focused on the vasculature, aiming to understand its structural characteristics, functions, interactions with surrounding tissues, and the mechanisms underlying vascular-related pathologies. However, advancing our understanding of vascular biology requires more complex and physiologically relevant models that integrate physical, chemical, and biological factors. Traditional *in vitro* dish models cannot replicate three-dimensional (3D) architecture, multi-cell-type interactions, and extracellular environments. *In vivo* animal models, while more complex, present ethical concerns, high costs, and limited relevance to human physiology. As a result, increasing attention is being directed toward *in vitro* models, specifically vascular microphysiological systems (MPS) based on organ-on-a-chip (OoC) technologies. This review highlights the relevance and potency of vascular MPS, which leverage microfluidic channels and 3D structures to mimic the physiological environment, incorporate diverse cellular and acellular components, and support complex biological processes. Vascular MPS are already enabling deep investigation into vascular responses to physiological cues, interactions with healthy and pathological tissues, and applications in disease modeling and drug development.

Received 6th January 2025,
Accepted 6th June 2025

DOI: 10.1039/d5lc00014a

rsc.li/loc

I. Introduction

Vasculature is one of the most critical systems for maintaining homeostasis, enabling blood circulation, delivering oxygen and nutrients to all organs, and removing waste for excretion. Given its essential role, dysregulation of the vasculature is related to numerous diseases and pathologies. Two major pathologies directly affecting vasculature are cardiovascular diseases, representing 37% of noncommunicable disease-related deaths in 2021, with roughly 20 million deaths globally,¹ and hypertension, affecting roughly 1.28 billion people worldwide.² In addition, non-vascular-specific diseases, such as various degenerative diseases³ and diabetes,⁴ either result from or contribute to vascular dysfunction. Tumor vasculature also plays a key role in cancer progression and is recognized as one of the hallmarks of cancer.^{5,6}

Traditional research models, including cell culture, animal models, and human studies, provided significant insights into vascular biology. However, each model has inherent limitations, including organizational relevance,^{7,8} proximity to human

physiology,^{9,10} sample availability,¹¹ or ethical concerns.^{10,11} Dish-based cell culture lacks physiological complexity, often relying on limited cell lines, facing challenges with multi-cell type co-culture and lacking 3D environment recapitulation. Animal models, while offering *in vivo* complexity, differ significantly from human physiology and raise both ethical and cost-related concerns. Human tissue studies, despite their physiological relevance and complexity, are constrained by limited availability, high sample to sample variability, ethical considerations, and lower experimental control.

In contrast, microphysiological systems (MPS) offer a promising alternative. These *in vitro* models incorporate diverse physiologically relevant cellular and acellular components, recapitulate 3D tissue architecture, and allow controlled manipulation of physical and chemical parameters. MPS are generally cost-effective, relatively user-friendly, and raise fewer ethical questions compared to traditional models. In recent years, the definition of MPS has come to encompass a broad range of technologies, including 3D cultures, 3D bioprinting, organoids, and organ-on-a-chip (OoC). Among these, MPS based on OoC technologies offer distinct advantages for vascular studies due to their microfluidic properties. Accordingly, this review uses the term “MPS” specifically to refer to OoC-based technologies, with other systems mentioned explicitly when relevant.

In this review, we highlight the current and recent advances in the use of MPS to model the vasculature, emphasizing their

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growing biological relevance. We first briefly introduce the vascular anatomy to underscore the complexity of *in vivo* vasculature and the need for appropriate models. This is followed by a description of the fabrication processes of polydimethylsiloxane (PDMS)-based MPS, particularly focusing on soft lithography. We then discuss how design strategies, both pre-patterning and self-organizing, enhance architectural fidelity. The physiological relevance of vascular MPS is further explored through considerations of cellular and acellular components. Finally, we highlight the power of vascular MPS in advancing our understanding of vascular mechanobiology, physiology, and pathophysiology.

I.1) Anatomy of the vascular system

The vascular system consists of blood vessels, looping from the heart to the different organs and back to the heart, and exhibits hierarchical organization, from arteries and arterioles to capillaries, then to venules and veins.

Arteries, arterioles, venules, and veins share a common structural organization, consisting of three distinct layers¹² (Fig. 1A). The innermost layer, tunica intima, comprises a continuous monolayer of endothelial cells (ECs) supported by a thin basement membrane. The middle layer is the tunica media, which is primarily composed of smooth muscle cells (SMCs) and has a vessel-type-dependent thickness. The outermost layer, the tunica externa, consists of fibroblasts embedded within an extracellular matrix (ECM).

Regarding the tunica media, arteries and arterioles are characterized by a thick layer, making them more contractile and elastic. SMCs in this layer regulate vessel stiffness through interaction with the ECM and are responsible for vascular contraction and dilation.¹³ In contrast, venules and veins have a thinner tunica media with limited contractile ability, relying instead on the contraction of surrounding muscles to facilitate blood flow.¹⁴ In the tunica externa, fibroblasts maintain the structural integrity and elasticity of the vascular wall by secreting ECM components, providing structural support, and anchoring the vessel to surrounding tissues.¹⁵ Despite this shared structural organization, vessel diameter and morphology vary widely. Arteries are generally round in cross-section, with elastic arteries ranging from 10 mm in diameter to 25 mm for the aorta, while muscular arteries range from 1 to 10 mm¹⁶ (Fig. 1A). Arterioles typically have diameters below 400 μm . Veins are more ovoid in shape, with diameters ranging between 5 mm and 15 mm, to 30 mm in the case of the vena cava¹⁶ (Fig. 1A).

Capillaries are composed solely of a monolayer of microvascular ECs, which can either be continuous, fenestrated, or discontinuous, depending on their function and anatomical location in the body.¹² Capillaries have diameters typically below 10 μm ,^{16,17} but form a dense vascular bed in direct contact with tissues. Their thin walls and extensive surface area make them the primary site of oxygen, nutrient, and waste exchange.¹⁷ The direct surrounding of capillaries is composed of pericytes lining the

vessels (Fig. 1A), which regulate the formation and maintenance of capillary networks, as well as blood flow.¹⁷

It is important to highlight the presence of specialized vascular structures within the vascular system, with the examples of the aortic arch and venous valves. The aortic arch is located in the initial segment of the aortic artery, between the ascending and descending aorta, and gives rise to three major arteries supplying the upper body.¹⁸ The aorta branch's primary functions are to redistribute blood flow between the aorta and the three arterial branches and to help regulate blood pressure through mechanosensory feedback mechanisms.¹⁸ In contrast, venous valves are a morphological feature specific to veins in the lower extremities. These valves consist of two leaflets (bicuspid valves) formed by protrusions of the tunica intima, reinforced by collagen and elastic fibers, and lined with endothelial cells.¹⁹ The main role is to prevent the backflow of blood back to the lower body, counteracting the effects of gravity and the lack of continuous muscular peristalsis.¹⁹

I.2) Vascular MPS fabrication methods

MPS can be fabricated using a variety of techniques. In recent years, 3D printing techniques using plastics or biopolymers have gained prominence and have been extensively reviewed.^{20–25} Some techniques are based on the direct printing of devices using UV light to cure a photosensitive polymer, such as stereolithography (SLA), digital light processing (DLP), or two-photon photopolymerization (TPP). Other direct-printing techniques rely on the extrusion and deposition of resins or gels, such as fused UV-direct ink writing (UV-DIW), fused deposition modeling (FDM), coaxial extrusion, and material jetting (MJ). Additionally, indirect 3D-printing is commonly used for 3D-printed molds and sacrificial templates.²⁴

Various types of polymers can be used in these fabrication methods. Plastic polymers, notably including PDMS, polystyrene (PS), polyethylene terephthalate (PET), and poly(methyl methacrylate) (PMMA), are favored for their ability to form rigid structures with relatively good resolution, transparency, and biocompatibility. Resins, like polyethylene glycol monomethacrylate (PEGMA), are also employed, despite limitations such as opacity and poor biocompatibility. Bioprinting, which uses naturally derived materials such as alginate, collagen, gelatin, or ECM, has become increasingly widespread in recent years.²³

This review, however, focuses primarily on PDMS-based devices fabricated using soft lithography, as it remains the most common method used for MPS fabrication, since the original lung-on-chip.²⁶ The soft lithography process begins with the design of a photomask using computer-aided design (CAD) software.²⁷ Then, UV light is projected through the photomask onto a silicon wafer coated with a thin layer of photosensitive resin. Depending on the type of photoresist used, the resin is either crosslinked (negative resist) or dissolved (positive resist) upon exposure to UV light. The

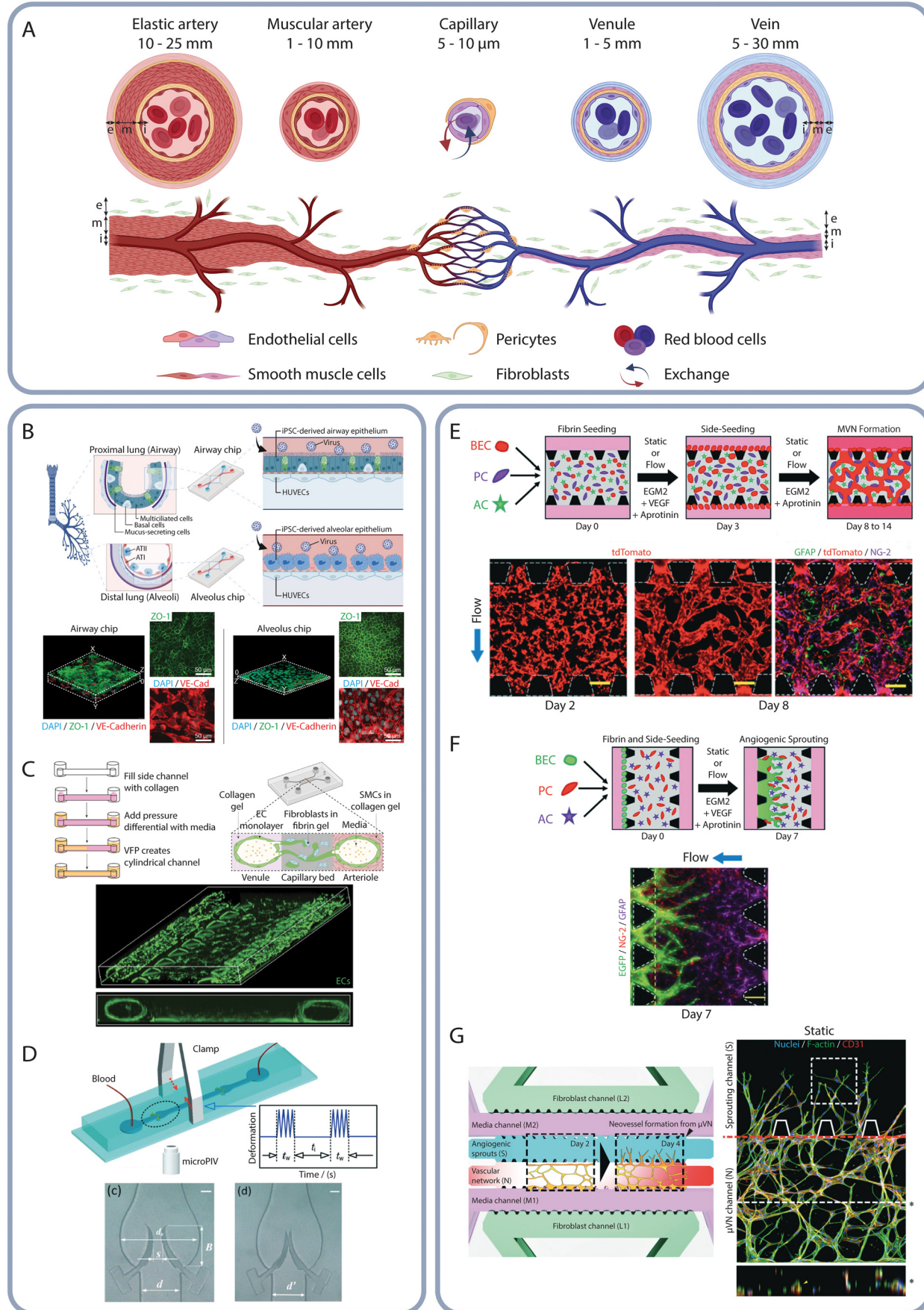


Fig. 1 Blood vessel architecture and diversity of vascular MPS models. A) The vascular system is composed of different-order vessels: arteries, arterioles, capillaries, venules, and veins. Vessels have specific structures and components, giving them diverse properties such as elasticity, permeability, and functions. Red blood cells flow through the vasculature. Gas, nutrient, and waste exchanges occur at the capillary-tissue interfaces. Tunica intima (i), tunica media (m), and tunica externa (e) are shown. Created in BioRender. Fujimoto, K. (2025) <https://BioRender.com/l33p826>. B-D) Pre-patterning methods to obtain vascular MPS. B) 2D bilayer devices modeling lung-on-chip.²⁹ The tissue channel is composed of either airways or alveolar epithelium, while the vascular channel is lined with ECs. SARS-CoV-2 infection has been recapitulated using these devices. Scale bars: 50 μ m. C) Using a viscous finger patterning method, 3D pre-patterned channels have been obtained and lined with ECs.³⁵ The figure has been rearranged from the original. D) A venous valve design, with two valves and a clamp in between, which mimics muscle contraction.⁴² (c) and (d) The open and closed state, respectively. t_i is the interval period with no clamping. t_w is the working period with ongoing clamping. d_s , S , d , d' , and B are various size parameters of the device. Scale bars: 100 μ m. E-G) Self-organizing methods are used to obtain 3D vascular networks *on-chip*, relying on either vasculogenesis or angiogenesis processes. E) A vasculogenesis-based method using a 3-channel device with micropillars.⁵³ Gel, containing a mixture of ECs, pericytes, and astrocytes, is injected into the central channel, and the vascular network forms spontaneously in 8 days. Scale bars: 200 μ m. The figure has been rearranged from the original. F) An angiogenesis-based method using the same device as E).⁵³ Gel, containing only pericytes and astrocytes, is injected into the central channel. ECs are seeded on the side of the gel and sprout over the course of 7 days. Scale bar: 200 μ m. The figure has been rearranged from the original. G) 6-channel device encapsulating both vasculogenesis and angiogenesis processes.⁵⁷ The vascular network forms during the first 2 days and sprouts start thereafter for 3 days. The figure has been rearranged from the original. All figures have been edited to enhance readability.

resulting master mold is often treated *via* siloxane-based coating to facilitate demolding. Liquid PDMS is then poured onto the master mold, degassed to remove air bubbles, and thermally cured. Finally, PDMS layers are cut, punctured, treated, and bonded to form the desired device structure.

Compared to other fabrication techniques, soft lithography enables high-resolution patterning. Qin *et al.* described the different resolutions in each step of soft lithography using a commercial printer.²⁷ Mask design can reach down to 1 μ m resolution, photomask printing down to 20 μ m, master fabrication down to 1 μ m, and PDMS stamping down to 500 nm. Even higher resolution can be achieved using chromium (Cr) photomasks with conventional photolithography. Nevertheless, typical soft lithography workflows can routinely produce devices with an approximate resolution of 20 μ m. Beyond resolution, PDMS offers several other advantages that make it a widespread material for MPS fabrication, including low cost, optical transparency, gas permeability, and biocompatibility.²⁸ However, notable drawbacks of PDMS include its inherent hydrophobicity and the tendency to absorb small molecules.²⁸ Owing to many favorable characteristics, PDMS has been extensively used in the field and remains the predominant material for MPS fabrication. This widespread adoption justifies the focus of the present review on PDMS-based systems.

II. Vascular MPS as highly relevant models

II.1) Design considerations for architectural relevance

Vascular MPS can be constructed using two methods: pre-patterning or self-organizing methods. While pre-patterning methods focus on either endothelium-tissue interfaces or blood vessel structure modeling, self-organizing methods aim to reproduce physiological phenomena *on-chip*.

Pre-patterning methods: 2D and 3D. In pre-patterning methods, cells adhere to the walls of pre-existing vessel structures, relying on microfabricated channel designs to define vessel diameter, length, connecting points, or overall

shape. The most traditional and widely used model, derived from Huh *et al.*'s original lung-on-chip,²⁶ is the two-dimensional (2D) bilayer structure. The designs consist of two superimposed channels separated by a porous membrane. A vascular side is lined with ECs and exposed to either static^{29,30} or flow^{31,32} conditions, while an organ side contains organ-specific cells, which can be epithelial cells,^{29,30} or perivascular cells^{31,32} (Fig. 1B). Variations in these 2D designs arise from differences in channel dimensions (e.g., length, width, height) and contact areas. These 2D vascular MPS are commonly used to study interactions between vasculature and tissues, offering valuable insights into both healthy^{26,32} and pathological^{29–31} conditions. However, although 2D bilayer models allow the application of physical cues such as flow or stretch, they share several limitations with culture on insets: namely, being restricted to bilayer co-culture and lacking 3D environments and structural complexity.

For this reason, pre-patterned 3D designs have recently gained momentum for their improved physiological relevance. However, constructing 3D pre-patterned models presents greater challenges, as traditional soft lithography techniques produce rectangular microfluidic channels that deviate from the circular geometry of *in vivo* blood vessels. While some studies turn to alternative fabrication methods such as 3D printing³³ and bioprinting,²¹ some have succeeded in obtaining 3D vessels even within rectangular channels using viscous finger patterning.^{34–36} This technique involves injecting gel into the channel and applying pressure or aspiration at one of the ends. Thus, removing the gel from the channel and leaving behind a tubular structure resembling the circular geometry of *in vivo* blood vessels. Chen *et al.* demonstrated the effectiveness of viscous finger patterning by creating a hierarchical microvasculature-on-chip that incorporated an arteriole surrounded by SMCs, capillaries, and a venule within a single device³⁵ (Fig. 1C). Simpler methods consist of seeding ECs on two or all walls to obtain lumen formation.^{37,38} Overall, these 3D models improve vessel architecture modeling, both in shape and

organization, with the integration of perivascular cells such as SMCs³⁵ or macrophages.³⁶

Pre-patterned devices have also been utilized to model specific vascular structures, such as the aortic branch and venous valves. For instance, Li *et al.* developed a device with an arched channel and three branches connected to the peak of the arch.³⁹ This model was used to assess shear stress distribution within the device and thrombosis conditions. In the case of venous valves, several studies have proposed variations of valve designs to investigate thrombosis.^{39–44} While most studies employed fixed valve structures, one specific study introduced small, flexible leaflets made of polyethylene glycol diacrylate (PEG-DA) that respond to flow direction, opening under forward flow and closing under backflow, thereby mimicking native valve behavior⁴² (Fig. 1D). Additionally, in this model, flow was controlled using a clamping mechanism on the side of the device that periodically compressed the channels to simulate calf muscle contractions. All together, these examples demonstrate how vascular MPS can be tailored to replicate complex anatomical features by integrating thoughtful microfluidic design, advanced fabrication processes, and functional mechanical components.

Self-organizing methods: vasculogenesis and angiogenesis. Self-organizing methods leverage biological processes occurring during vascular development, usually during prenatal stages, including vasculogenesis and angiogenesis. Vasculogenesis refers to the *de novo* formation of a vascular network from a pool of endothelial progenitor cells present in the tissue, while angiogenesis involves the sprouting of new vessels from pre-existing vessels through EC migration.⁴⁵ Another related process, known as neovasculogenesis, occurs postnatally, particularly during tissue repairs or pathological conditions.⁴⁶ However, neovasculogenesis has not yet been applied in vascular MPS. This process describes the formation of new vessels within an already existing vascular network, typically originating from circulating endothelial progenitor cells.⁴⁶

Vascular MPS based on vasculogenesis consists of ECs embedded within a hydrogel, where they spontaneously form vascular networks over time (Fig. 1E and G). Common designs include 3-,^{47–53} 5-,^{54–56} or even 6-channel⁵⁷ configurations, separated by micropillars. The central region, which houses the vascular network, can take various shapes, such as diamond-shaped^{47–49} or rectangular.^{51,53,54,57} Side channels serve as medium reservoirs, perfusion channels or may also be embedded with other cell types. Vasculogenesis is most often induced by growth factors secreted by fibroblasts, which are either co-embedded with the ECs⁴⁹ or placed in the side channels.^{54,55,57} In some studies, cancer cells have been used to initiate vasculogenesis.^{48,50} Similarly, fibroblast-free vascular MPS have also been developed, using growth factor-rich media containing vascular endothelial growth factor (VEGF) and fibroblast growth factor (FGF)^{47,51–53,56} to induce network formation. These vasculogenesis-based models are particularly valuable for studying interactions between ECs and organ-specific, tumor, or perivascular cells

through direct co-culture within the gel^{51,53} or by culturing them on top of the vascular network.^{55,56}

Angiogenesis-based vascular MPS designs are often similar or even identical to those used in vasculogenesis-based models. However, in angiogenesis models, the gel-containing region is either lined with ECs that will sprout into the gel⁵³ or contains a pre-existing vascular network from which sprouts emerge⁵⁷ (Fig. 1F and G). Sprouting is stimulated by specific signals such as growth factors,^{53,58} fibroblasts,^{51,54,55,57,59} or cancer cells⁶⁰ embedded within or across the gel. These models are particularly useful to study angiogenic potential in the presence of perivascular cells^{51,53} or tumor cells,⁶⁰ or the response to physical cues⁵⁷ or therapeutic drugs.⁵⁷

In contrast to vasculogenesis and angiogenesis, which are well-characterized and widely used in vascular MPS, neovasculogenesis remains relatively poorly understood and has not yet been implemented in the field. Only a limited number of studies have explored this process, showing its critical roles in tissue repair and pathological conditions.⁶¹ Therefore, the development of vascular MPS models capable of recapitulating neovasculogenesis would provide valuable insights into its underlying mechanisms and functional relevance.

II.2) Cellular and acellular considerations for physiological relevance

In vascular MPS, the EC source is the most critical factor for achieving physiological relevance, as ECs are the cornerstone of vasculature. Perivascular cells, such as fibroblasts, pericytes, and SMCs, are equally important since blood vessels *in vivo* are composed of multiple cell types, required for their structure, function, and stability. In addition to cellular components, the microenvironment plays a major role in vascular physiology and pathology. Key elements include the basement membrane, which can be mimicked with artificial membranes, and the ECM, which is recreated using hydrogels. Together, these components create a more realistic environment that supports vascular formation, function, and the study of disease models.

Endothelial cells: umbilical, cell lines, organ-specific, and progenitors. The source of ECs is a critical factor in constructing *in vivo*-like vasculature and vascularized tissues or organs. Human umbilical vein ECs (HUVECs) are conventionally used in MPS research, both in pre-patterning and self-organizing methods (Fig. 2A), as they are easy to source and culture, and most importantly for their ability to form vascular structure and express genes associated with EC phenotype, notably tight junctions.^{54,62–64} HUVECs serve as a versatile tool facilitating studies on vascular development, barrier functions, and responses to chemical stimuli.^{58,65,66} In pre-patterning methods, HUVECs are mainly used to investigate barrier functions in interaction with epithelial cells.^{29,64,67} In self-organizing methods, they enable the analysis of vasculature morphology over time, including

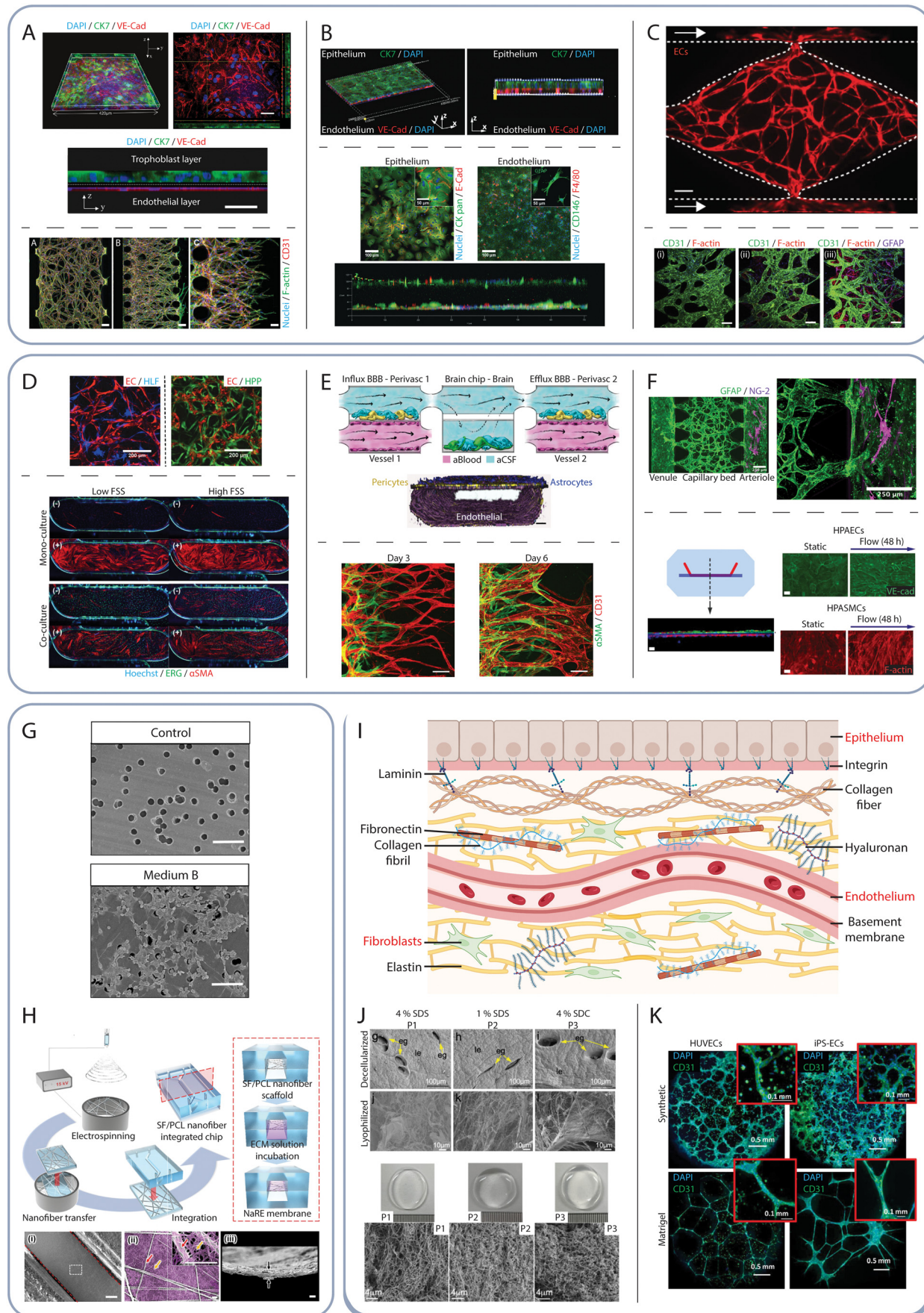


Fig. 2 Diversity of ECs and perivascular cells, and importance of extracellular components: membrane and matrix in vascular MPS. A) HUVECs are the most commonly used EC type in vascular MPS. Top panel shows a 2D bilayer device composed of a trophoblast epithelium and HUVECs, recapitulating a placental model.⁶⁷ Scale bars: 50 μm and 30 μm . The figure has been rearranged from the original. Bottom panel represents both vasculogenesis (A) and angiogenesis (B and C) methods to form a 3D vascular network with HUVECs.⁵⁴ Scale bars: 100 μm (A and B) and 50 μm (C). The figure has been rearranged from the original. B) Organ-specific ECs in 2D bilayer models. Top panel illustrates a lung-on-chip model with alveolar epithelial cells (HPAEpiC) and human lung microvascular ECs (HULEC-5a line).³⁰ Bottom panel displays a liver-on-chip model, encapsulating hepatocytes (left and inset) liver sinusoidal ECs (right), and hepatic stellate cells (right inset) isolated from murine liver samples.⁸² Scale bars: 100 μm and 50 μm . The figure has been rearranged from the original. C) Progenitor-derived ECs in 3D vascular MPS. Top panel presents a 3D vascular network derived from ECFC-ECs.⁴⁸ Scale bar: 100 μm . Bottom panel is a 3D BBB-on-chip model created using iPSC-ECs monoculture (i), or in co-culture with brain pericytes (ii), or both pericytes and astrocytes (iii).²⁰⁸ Scale bars: 100 μm . D) Fibroblasts are commonly used as a tool, but rarely are the focus of studies. Top panel shows a 3D HUVEC-based vascular network in contact with human lung fibroblasts (HLF in blue) and human placental pericytes (HPP in green).⁶⁸ Scale bars: 200 μm . Bottom panel is a 2D bilayer device with primary human retinal microvascular ECs and dermal fibroblasts, in the presence (+) or absence (-) of 10 ng mL^{-1} TGF β 1.⁹⁷ Channel size: 4 mm \times 1 mm. E) Pericytes are commonly used in contact with ECs. Top panel illustrates a 3-device BBB-on-chip model with brain pericytes.³² Scale bar: 75 μm . The figure has been rearranged from the original. Bottom panel is a 3D angiogenesis model with dermal pericytes lined to the vessels.⁵⁹ Scale bars: 100 μm . The figure has been rearranged from the original. F) SMCs are used to recreate artery or arteriole models. Top panel presents a 3D hierarchical vascular system, encapsulating a venule, capillaries, and an SMC-lined arteriole.³⁵ Scale bars: 250 μm . The figure has been rearranged from the original. Bottom panel displays a 2D bilayer pulmonary artery-on-chip with pulmonary arterial ECs and pulmonary arterial SMCs, under flow conditions.³¹ ECs align parallel to flow, while SMCs align perpendicularly. Scale bars: 10 μm . The figure has been rearranged from the original. G) Scanning electron microscopy (SEM) images from the Transwell membrane without or with coating, composed of fibrinogen and growth factors (medium B).²⁸⁵ Scale bars: 50 μm . The figure has been rearranged from the original. H) Protocol for the formation of electrospun membranes, composed of SF-PCL fibers and ECM gel.¹¹⁹ SEM images of the membrane (i, ii, and inset). The red arrow highlights SF-PCL fibers and yellow collagen nanofibrils. iii shows a cross section image of the membrane. Scale bars: 500 μm and 2 μm . I) Schematic of ECM organization. Red words are cellular components, black words are ECM components. Created in BioRender. Fujimoto, K. (2025) <https://BioRender.com/y31y833>. J) SEM images of decellularized (g-i) and lyophilized (j-l) endometrium tissues, following three protocols (P1-3).¹⁵⁷ Scale bars: 100 μm and 10 μm . Gross and SEM pictures of the dECM-derived hydrogels at a concentration of 10 mg mL^{-1} . Scale bars: 4 μm . The figure has been rearranged from the original. K) Vascular network derived from HUVECs and iPSC-ECs, out-of-device, using Matrigel or synthetic PEG-based hydrogels.¹⁶⁶ Insets show magnified images of each gel. Scale bars: 0.5 mm and 0.1 mm. All figures have been edited to enhance readability.

parameters such as vascular diameter, connectivity, and length.^{54,58,68} Moreover, HUVEC-based self-organizing methods have been used to replicate *in vivo*-like barrier functions and to study drug effects.^{66,69} A less commonly utilized EC source is human umbilical artery ECs (HUAECs).⁷⁰ Combining HUVECs and HUAECs to create vascular networks allows researchers to model heterogeneous blood vessels, offering new insights into the roles of different EC types in vasculogenic processes.⁷⁰

Similarly, other types of ECs, including EC cell lines and immortalized ECs, have also been used in vascular MPS. Several studies have demonstrated their applicability for modeling vascular functions. For instance, the EC line EA.hy926 was used to construct a liver-on-chip model, in which hepatic functions were successfully maintained under flow conditions.⁷¹ Another example is the use of TeloHAECS, an immortalized human aortic EC line, in a pre-patterned vascular lumen model to evaluate tumor cell intravasations.⁷² Both studies highlight the feasibility of incorporating EC cell lines and immortalized ECs into vascular MPS. Overall, these different types of ECs already grant access to meaningful information about vascular development, barrier functions, and interactions with the environment.

However, using non-organ-specific ECs presents some limitations in recreating organ specificity or diseases *in vitro*. Both HUVECs and HUAECs are differentiated cells that express specific phenotypes associated with the umbilical cord.⁷³ For cell lines and immortalized cells, altered biology and genetic anomalies are often present, compared to *in vivo* cells. Meanwhile, it is well known that ECs have structural specificity and differences among organs: continuous,

fenestrated, and discontinuous ECs.⁷⁴ More recent advancements in multi-omics and single-cell RNA sequencing (scRNA-seq) technologies have further revealed significant EC diversity in humans and mice.^{75,76} This diversity also extends to pathological responses, as specific EC types react differently to disease conditions. For example, a recent study shows that kidney-derived ECs are more susceptible to diabetes-associated damage than ECs from other organs.⁷⁷ These findings emphasize the importance of using organ-specific ECs for constructing vascular MPS. Incorporating organ-specific ECs would enhance physiological relevance, enabling more accurate modeling of both healthy and pathological vascular environments.

Organ-specific microvascular ECs have been used in MPS to model tissue–capillary interactions and investigate organ-specific functions such as barrier integrity and molecular transport, particularly through pre-patterning methods. Continuous ECs, such as those present in the lungs and brain, are crucial for preventing harmful chemicals or pathogens from penetrating blood or tissue. The first MPS, a lung-on-chip, incorporated pulmonary microvascular ECs (HPMECs) to replicate a continuous phenotype.²⁶ Numerous other studies use different cell lines of human lung microvascular ECs^{30,78–80} (Fig. 2B top). Similarly, for brain models, human brain microvascular ECs (HBMECs) have been used to create a continuous and tight blood–brain barrier (BBB) in contact with pericytes and astrocytes, mimicking the *in vivo* environment.³² In contrast, fenestrated endothelium, essential for selective filtration, is characteristic of organs such as the kidney and small intestine. A model mimicking kidney fenestrated endothelium, using glomerular

microvascular endothelial cells (GMECs), successfully demonstrated caveolae-mediated transendothelial transport, replicating key filtration functions of the glomerulus.⁸¹ Finally, discontinuous endothelium, as found in the liver, facilitates efficient metabolic exchanges. A murine liver-on-chip incorporated liver sinusoidal ECs (LSECs) to recreate this specialized endothelium⁸² (Fig. 2B bottom). Additional examples of EC types and their applications in organ-specific MPS are reviewed by Urbanczyk *et al.*⁸³

While MPS using pre-patterning methods effectively recapitulate capillary diversity using organ-specific ECs, self-organizing methods have yet to achieve comparable versatility. Among MPS using self-organizing methods, BBB-on-chip has been well explored using brain-specific ECs, with, for instance, Winkelman *et al.* employing primary HBMECs to generate a 3D vessel network.⁵³ HBMECs display higher mRNA expression of permeability-related genes⁵¹ and better mimic *in vivo*-like permeability compared to HUVECs.⁸⁴ Other examples include human dermal microvascular ECs (HDMECs),⁵⁵ human colonic microvascular ECs (HCoMECs),⁸⁵ and HPMECs.⁸⁶ Despite these promising examples, the number of MPS capable of recapitulating 3D, organ-specific microvascular networks remains limited, highlighting a significant gap in modeling the structural and functional diversity of tissue-specific capillaries.

Alternative sources of ECs are progenitor cells and stem cells. Endothelial colony-forming cells derived ECs (ECFC-ECs) and induced pluripotent stem cell-derived ECs (iPSC-ECs) are particularly noteworthy. ECFC-ECs, which are resident cells within the vasculature, can migrate to injury sites and differentiate into specific EC types. They exhibit prolonged replication capacity^{87,88} and greater angiogenic potential than HUVECs,⁸⁹ making them ideal candidates for self-organizing methods (Fig. 2C top). Similarly, iPSC-ECs can form a perfusable vasculature *via* self-organization and differentiate into organ-specific ECs^{51,90} (Fig. 2C bottom). Moreover, iPSC-ECs offer the unique advantage of being sourced from patients with specific diseases, allowing the recapitulation of pathological development and phenotypes.^{52,91} This capability makes iPSC-ECs invaluable for creating both healthy and diseased models, advancing research into vascular biology, and opening new opportunities in personalized medicine.

Overall, the choice of EC source is a critical parameter in vasculature studies using vascular MPS. While HUVECs remain widely used in MPS, the field is increasingly adopting alternative cell sources to enhance physiological relevance. Organ-specific ECs and ECFC-ECs both offer substantial potential to enhance the fidelity of vascular models in MPS research. In the context of personalized medicine, iPSCs-ECs and patient-derived ECs represent promising approaches, enabling the generation of personalized, organ-specific vascular networks. One example, explored by Perry *et al.*, utilized patient-derived ECs from the limb veins in an out-of-device model to successfully form a 3D vascular network.⁹²

Perivascular cells: fibroblasts, pericytes, and smooth muscle cells. While ECs are the primary component of blood

vessels, perivascular cells, such as stromal cells (e.g., fibroblasts) and mural cells (e.g., pericytes and SMCs), are essential for proper vasculature organization and functions.^{74,93} Incorporating these supporting cells into MPS is a critical step toward developing more accurate and physiologically relevant models.

In vivo, fibroblasts reside within the ECM, playing a key role in ECM remodeling⁹³ and growth factor secretion.^{93–95} Consequently, fibroblasts are frequently used in self-organizing methods to support EC vasculogenesis or angiogenesis (Fig. 2D top), limiting fibroblasts as a simple tool for vasculature formation and overlooking their broader contributions to vascular biology. Meanwhile, crosstalk between ECs and fibroblasts is pivotal in both physiological and pathological contexts. For instance, a recent study demonstrated that fibroblasts can generate physical cues, such as ECM stiffening, which promote vasculogenesis and enhance blood vessel integrity.⁹⁶ Another study showed that ECs can alleviate myofibroblast activation mediated by TGF β signaling, highlighting crosstalk between these cell types⁹⁷ (Fig. 2D bottom). These studies emphasize the importance of exploring fibroblast-EC interactions in MPS, particularly under both normal and pathological conditions. By addressing these interactions, MPS can uncover insights from vascular biology and improve the functionality of engineered vascular MPS.

Mural cells, another type of perivascular cell, are in direct contact with vessels.⁹³ These include pericytes, which are close to capillaries, and SMCs, which are associated with arteries and veins.⁷⁴ Both cell types play crucial roles in maintaining vascular integrity and stability.⁹⁸ Pericytes are usually incorporated into MPS to replicate tissue-specific vascular characteristics, especially for lung or brain modeling. As pericytes are critical for maintaining the BBB, numerous studies have encapsulated pericytes *on-chip*^{32,51,59} (Fig. 2E). Pericytes allow the recapitulation of capillary-like phenotypes by enabling the formation of smaller-diameter vessels in self-organizing methods.^{68,88} Additionally, encapsulating pericytes within vascular MPS allows the modeling of another hallmark of lung or brain capillaries: their reduced permeability. MPS incorporating lung or brain pericytes have successfully achieved this characteristic, further enhancing physiological relevance.^{32,59}

SMCs, on the other hand, are used to recapitulate artery-like blood vessels.^{31,35} In a pre-patterning method, SMCs can be incorporated into relatively large 3D blood vessels to achieve an *in vivo*-like environment (Fig. 2F top).³⁵ Another notable example is the development of a three-layer arteriole model, consisting of ECs, SMCs, and fibroblasts, representing the tunica intima, media, and externa, respectively.⁹⁹ The authors successfully replicated early-stage atherosclerosis by applying various initiating stimuli, such as low-density lipoprotein (LDL) and tumor necrosis factor α (TNF α). The authors investigated vasoactivity, permeability, and monocyte phenotypes, as well as evaluated the efficacy of therapeutic drugs. In a self-organizing model, iPSC-derived SMCs have been utilized to investigate the

interactions between SMCs and vascular structures, focusing on the relationship between SMC spatial localization and contractile function.¹⁰⁰ Alternatively, in more traditional 2D bilayer devices, SMCs are employed to study diseases affecting arterial walls or SMC-specific functions, such as pulmonary arterial hypertension (PAH) (Fig. 2F bottom).³¹ Although some vascular MPS incorporating SMCs are already established, further refinement and expanded investigation are warranted, given the critical role of SMCs in vascular physiology and disease.

Like ECs, perivascular cells present heterogeneous and organ-specific gene and protein expression patterns.¹⁰¹ Advances in multi-omics technologies revealed this diversity both between and within organs.^{101–104} Fibroblasts demonstrate inter- and intra-organ heterogeneity across various tissues, including the heart, skeletal muscles, colon, bladder,¹⁰¹ brain,¹⁰² liver,¹⁰³ and kidney.¹⁰⁴ Therefore, fibroblasts from different organs are likely to influence vascular morphology differently, as seen with lung fibroblasts producing larger diameters compared to dermal fibroblasts or bone marrow-derived mesenchymal stem cells.¹⁰⁵ Pericytes have also exhibited organ-specific functionality in vascular MPS. Patient-derived CD90+ CD146+ pericyte-like cells from lung tissues have been used to develop chronic obstructive pulmonary disease *on-chip*.¹⁰⁶ Similarly, both primary and iPSC-derived brain pericytes have been successfully used to construct the BBB,^{32,107} highlighting the value of organ-specific pericytes in generating physiologically relevant MPS. Additionally, non-perivascular cells have also been incorporated in vascular MPS, such as astrocytes for brain,^{32,51} stellate cells for liver,⁸² and mesangial cells for kidney glomeruli.¹⁰⁸ Overall, similarly to ECs, using organ-specific perivascular and non-perivascular cells in vascular MPS is key for improved relevance.

Membrane: current and future generations. In 2D bilayer MPS, membranes are key in separating compartments and supporting cell growth. These membranes are designed to mimic the basement membrane found *in vivo*, which separates endothelium from surrounding tissues.

The most widely used membranes are PET porous membranes (Fig. 2G), with pore sizes ranging from 0.4 μm to 8 μm and a thickness of approximately 8–10 μm , making them suitable for a wide range of applications, such as brain,^{32,109} lung,^{29,31} or placenta models.⁶⁷ Other materials are also widely used; for instance, PDMS can be spun into thin layers to create membranes. Notable examples include a 10 μm -thick PDMS membrane used in the original lung-on-chip.²⁶ Another study uses 25 μm membranes with 5 μm pores for a gut-on-chip model.¹¹⁰ Polycarbonate (PC) is another frequently used material, offering membranes with a thickness of around 10 μm and pore sizes ranging from 0.4 to 10 μm .^{32,111,112} More recently, silicon nitride (SiN) membranes have gained attention in applications such as BBB-on-chip to study particle translocation and transcytosis.^{113,114} SiN membranes are exceptionally thin, with a thickness of approximately 100 nm, closer to the <100 nm of the *in vivo* basement membrane.^{115,116} These

membranes are highly permeable, capable of sustaining cell culture, and provide superior imaging resolution compared to traditional materials.^{113,114}

The membranes previously described present two shortcomings: a higher thickness and a less relevant structural organization compared to *in vivo* conditions. These membranes are typically flat, relatively smooth, and less elastic, whereas the natural basement membrane comprises collagen and other proteins arranged into fibrous, mesh-like sheets that provide elasticity and structural complexity.^{116–118} To overcome such shortcomings, electrospun membranes have emerged as a promising alternative. They are thinner (<5 μm), highly permeable, and composed of fibers resembling *in vivo* bundles with diameters of <100 nm.¹¹⁹ Electrospun membranes are increasingly used in MPS, particularly for modeling organs subjected to mechanical cues, flow, or stretch. Examples include MPS with integrated vasculature^{119,120} (Fig. 2H) and those without vasculature,^{121–123} both of which show the elastic properties of these membranes. Broadly speaking, materials range from plastic polymers such as poly(methyl methacrylate) (PMMA)¹²¹ or polycaprolactone (PCL)^{119,120,122,123} to biopolymers such as silk,^{119,120,124,125} collagen,^{123,126} and composites like chitosan.¹²⁷ However, in vascular MPS, the combination of both plastic- and biopolymer-based membranes shows great promise, as highlighted in Kanabekova *et al.*,¹²³ where they use PCL and collagen.

Finally, despite some examples of biopolymer-based membranes, most membranes used in vascular MPS are made of plastic or other synthetic materials. While these membranes are generally biocompatible, their intrinsic material properties continue to limit their ability to fully replicate physiological environments. Key characteristics, including elasticity, permeability, and cell adhesion capability, still fall short when compared to the native basement membrane. Therefore, further advancements in membrane materials and fabrication techniques are necessary to enhance their functional resemblance to *in vivo* conditions and support more physiologically relevant tissue models.

Currently, to compensate for the cell binding capability, numerous studies use a subsequent coating, notable to facilitate cell adhesion and growth, mimicking the composition of basement membrane proteins. The choice of coating agents is diverse and depends heavily on the cell type being used, requiring optimization to find the most suitable match. Collagen^{26,31,32,67,110,112–114,119} and fibronectin^{26,32,109,113,114} are the two most commonly used coating agents (Fig. 2G). In some cases, specialized coatings are employed to cater to specific cell types. For example, Yadav *et al.* used iMatrix-511 for airway chips and Geltrex for alveoli chips, to match the sensitivity and specificity of each iPSC-derived cell.²⁹ Typically, the choice of coating agent is guided by a combination of trial-and-error and previous literature, making it a case-dependent parameter.

Matrix: convention, alternative, and perspective. In 3D devices, the ECM plays a major role in device design for vessel development. The vascular ECM is a complex network of

proteins and carbohydrates, providing structural support and regulating cellular functions *via* signaling molecules, such as cytokines and growth factors.¹²⁸ The ECM also supports the formation and maintenance of vascular networks¹²⁹ by facilitating EC proliferation, migration, and adhesion, through biochemical and mechanical cues.¹³⁰ Additionally, the vascular ECM is involved in various pathologies, most notably fibrosis and cancer.¹³¹ These diverse properties make the ECM a crucial component for developing physiologically and pathologically relevant vascular MPS.

In vivo vascular ECM encapsulates multiple components, such as proteins, glycoproteins, and polysaccharides (Fig. 2I). Proteins include collagen, the most abundant and ubiquitous component, which provides structural support and tensile strength to blood vessels;¹³² elastin, which is responsible for elasticity, particularly in artery walls;^{132,133} and laminins, which are important for the structure and function of the basement membrane.^{132,134} Glycoproteins include proteoglycans, which maintain structural integrity and regulate the availability of growth factors, cytokines, and water,^{135,136} and fibronectin, which serves as a critical linker between the ECM and cells.^{137,138} Lastly, glycosaminoglycans, such as hyaluronic acid, are the main polysaccharides contributing to the ECM's hydration and structural properties.^{135,139} However, due to technical limitations, replicating the exact composition of the *in vivo* ECM remains challenging *in vitro*. In vascular MPS, the ECM has been widely mimicked using fibrinogen and collagen, with additional alternatives such as gelatin, alginate, and Matrigel being widely employed.

Fibrinogen is one of the most used ECM components in vascular MPS.^{51,52,62,68,70,84,87,88} The standard protocol requires fibrinogen to be cleaved into fibrin using thrombin, which then polymerizes into a gel to support lined or embedded cells. Despite fibrinogen being prevalent, collagen type I remains another frequently employed ECM component in MPS.^{63,90,140,141} Collagen protocols typically require pH neutralization and temperature-dependent polymerization to generate stable gels. Some studies have combined both fibrinogen and collagen to leverage their complementary properties.^{54,65,142,143} The final concentration of each component is case-dependent and influenced by various parameters such as cell type, vascularization method, and experimental outcomes. For fibrinogen, concentrations usually range between 2 and 10 mg mL⁻¹, while collagen concentrations vary between 2 and 6 mg mL⁻¹ when used alone and around 0.2 mg mL⁻¹ when mixed with fibrinogen. Helm *et al.* showed in an out-of-device model that both fibrinogen and collagen concentrations impacted vascular network development.¹⁴⁴ Similarly, Whisler *et al.* showed in an *on-chip* study that increasing fibrinogen concentration enhanced network branching, with the branch surface area increasing proportionally.⁶² Overall, fibrinogen and collagen are the most widely used hydrogels to mimic the ECM in vascular MPS, despite their relatively limited relevance.

To achieve greater physiological relevance, animal-derived gels are also employed. Gelatin, a collagen-rich material derived from animal tissues, can be used alone or in combination with other components, such as polyethylene glycol (PEG) or methacrylic anhydride (MA). For instance, gelatin-MA has been used to embed cells in a 2D bilayer device,¹⁴⁵ while gelatin with or without PEG has been used to generate a lumen scaffold in a 3D pre-patterning method.¹⁴⁶ Another common option is Matrigel, a native basement membrane matrix derived from the Engelbreth-Holm-Swarm mouse tumor. Matrigel is rich in laminin, collagen IV, and growth factors, making it highly effective in promoting angiogenesis and vascularization.¹⁴⁷ While Matrigel is commonly used for organoid and spheroid formation, Matrigel has also been successfully applied in MPS studies.¹⁴⁸⁻¹⁵⁰

One of the most promising alternatives is decellularized matrix (dECM). Derived from native tissues, dECM can originate from various species (human, rat, mouse, pig) and organs (lung, brain, liver, kidney). Decellularization processes remove cellular components while preserving the ECM components and structures.¹⁵¹⁻¹⁵³ This is achieved using a combination of chemical methods (detergents such as Triton X-100, SDS, hyper-/hypotonic solutions), enzymatic methods (trypsin or nuclease), and physical methods (freeze-thaw cycles, sonication, scraping, or ultra-high pressure). Most protocols combine these approaches to ensure effective cell removal while maintaining ECM integrity. Following decellularization, dECM can be used as a scaffold for tissue culture and engineering,¹⁵¹ or reduced into powder form to create hydrogels.¹⁵²⁻¹⁵⁴ Subsequent hydrogels have been applied in cancer modeling,¹⁵⁴ organoid culture^{153,155-159} (Fig. 2J), and MPS.¹⁶⁰⁻¹⁶⁴ In MPS, dECM has primarily been utilized to study organoid development,¹⁵⁵ skin aging,¹⁶² or anti-cancer drug efficiency,^{160,163,164} but has not yet been adopted for vascular studies. Despite the availability of commercially produced dECM,¹⁶⁵ its broader application remains limited by challenges related to ethical considerations and sample availability, as dECM is derived from animal or human tissues. For this reason, its use in vascular MPS is currently limited. However, several avenues exist for its application in vascular MPS, such as testing commercially available materials, fostering collaborations between researchers and clinicians to source tissue ethically, or leveraging future advancements in the dECM field. Nevertheless, due to its high physiological relevance and versatility, dECM holds strong potential as a biomimetic material for vascular research in MPS.

Lastly, synthetic gels, including PEG and polyacrylamide, have gained attention as alternatives to natural gels. These materials offer advantages such as well-defined composition, reproducibility, and stable mechanical and biochemical properties. Nguyen *et al.* highlighted the superior consistency of PEG compared to Matrigel, showcasing its potential for EC culture and vascular modeling, in an *out-of-chip* model¹⁶⁶

(Fig. 2K). Additionally, HUVECs have been shown to form vascular networks within a PEG-containing device.¹⁶⁷

III. Vascular MPS as potent tools

III.1) Recapitulation and study of vascular physical aspects

Vascular MPS enable precise reproduction and investigation of various physical parameters that influence vascular function. *In vivo*, blood vessels are constantly exposed to mechanical forces arising from hemodynamic activity, such as shear stress and interstitial flow. These forces are essential not only for vascular development and homeostasis but also play significant roles in the progression of vascular pathologies. Furthermore, the vasculature is responsible for the exchange of gases, nutrients, and waste products, while simultaneously acting as a selective barrier that prevents the infiltration of harmful substances. This selective transport is governed by endothelial permeability, another crucial parameter that can be studied using vascular MPS.

Luminal flow: shear stress or static. Fluid shear stress is one of the most extensively studied mechanical cues in vascular MPS. *In vivo* experiments have been implemented and revealed a strong coupling between shear stress and physiological or pathological processes, such as atherosclerosis or atheroma formation.^{168,169} Leveraging these insights, vascular MPS has been intensively used to study shear stress, thanks to the inherent channel-based fluidic designs. By controlling geometry and flow rate, vascular MPS can replicate *in vivo*-like conditions with high fidelity.

Shear stress can be applied across various MPS types, including 2D bilayer devices,^{31,170–172} 3D models based on pre-patterning methods,¹⁷³ and self-organizing 3D vascular networks.¹⁴² In both 2D and 3D systems, exposure to shear stress induces alignment of ECs to the flow direction, both in their overall orientation and cytoskeleton organization, mimicking *in vivo* EC behavior in blood vessels^{174,175} (Fig. 3A). SMCs also respond to shear stress by aligning perpendicularly to the direction of flow.³¹

In 3D self-organizing vascular MPS, the effect of flow and shear stress on angiogenic sprouting remains controversial.^{141,173,175,176} Some studies show that shear stress inhibits angiogenesis^{141,175} (Fig. 3A), whereas another study suggests that angiogenesis may be triggered only after a specific shear stress threshold.¹⁷³ A possible explanation, proposed by Wragg *et al.*,¹⁷⁶ is that angiogenesis is not triggered by shear stress itself, but rather by changes in shear stress within the vascular network. The timing of flow initiation and the magnitude of variation between baseline and flow conditions may, therefore, result in divergent outcomes. These contradictory findings emphasize the need for further exploration of shear stress and its relationship with angiogenesis. When it comes to vasculogenesis, shear stress has also been associated with the enhancement of network expansion, as evidenced by an increased number of branches, junctions, and endpoints.¹⁴² Shear stress has also been shown

to modulate mRNA and protein expression in ECs, including upregulation of endothelial nitric oxide synthesis (eNOS), a hallmark of EC function highly linked to shear stress,¹⁷⁶ as well as other markers such as thrombomodulin¹⁷⁰ and angiotensin-converting enzyme 2 (ACE2).¹⁷⁷

Despite its importance, many MPS studies overlook the incorporation of flow and shear stress, often opting for static conditions. This may be attributed to a lack of reference values for *in vivo* blood flow rates and shear stresses, as well as the challenges associated with quantifying shear stress within devices. Computational simulations can be used to address these challenges, either by determining the precise flow rates required to achieve a target shear stress^{31,175} (Fig. 3B) or by retrospectively analyzing the flow conditions applied.¹⁴² Real-time measurements in microfluidic channels can also help evaluate the flow rate and shear stress more accurately.¹⁷⁸

Overall, the development of flow generation systems has further expanded the applicability of shear stress in MPS. To date, syringe pumps^{142,173,175} and peristatic pumps^{31,170,174,175} have been commonly utilized to generate flow. These devices are relatively affordable and readily accessible; however, they often require complex experimental setups involving extensive tubing and bubble traps. Alternative methods, such as pressure-driven flow or magnetic stirrers, have also been used.¹⁷⁹ While these methods are simpler to implement and require less intricate setups, they usually offer limited control over the flow rate and may struggle to maintain stable, continuous flow. Improvements have, however, been achieved in a pressure-driven method by incorporating pumps and pressure sensors to maintain a constant pressure difference.¹⁸⁰ More recently, applications of oscillatory flow systems using a piezoelectric pump¹⁷¹ or pneumatic pressure¹⁷² showcase the potential for more precise and reproducible control of fluidic conditions. These systems offer improved precision and reproducibility in controlling fluid dynamics within MPS. Given the importance of luminal flow and shear stress in vascular biology, continued development of innovative flow generation strategies is essential to facilitate more physiologically relevant studies and to fully realize the potential of vascular MPS.

Other flows: interstitial and transluminal flow. Interstitial flow (IF) refers to the slow, pressure-driven movement of fluids through the ECM and interstitial spaces surrounding blood vessels. IF facilitates the transport of nutrients, waste, and signaling molecules to peripheral cells. Beyond its transport role, IF is now known to play a crucial role in vascular morphogenesis, including tumor angiogenesis, by influencing the distribution and activity of growth factors.¹⁸¹ To mimic the intricate physical environment associated with IF, 3D MPS models have been used to apply flow within channels and across the ECM, notably using hydrostatic pressure¹⁸² (Fig. 3C).

One of the key findings from MPS studies is that IF promotes angiogenesis,^{53,57,183} with a notable sensitivity to the flow direction (Fig. 3D). For instance, vascular sprouting is enhanced when flow is directed from the ECM towards the vessel.⁵⁷ More recently, the role of IF on vasculogenesis and

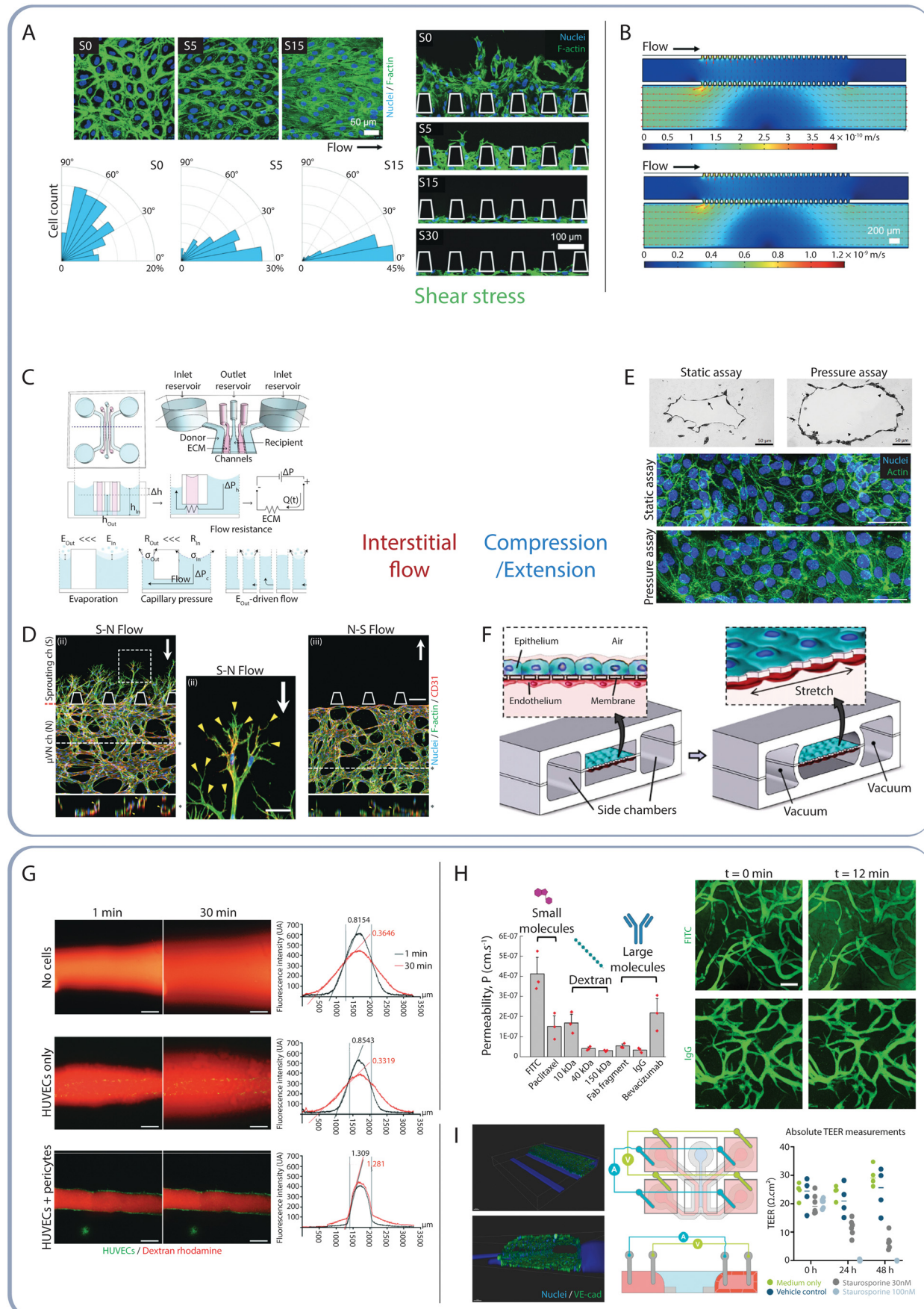


Fig. 3 Physical aspects in vascular MPS, to study flow, stresses and permeability. A and B) Studies of shear stress in vascular MPS. A) Effect of flow and shear stress on the EC monolayer and angiogenic sprouts.¹⁷⁵ Both EC monolayer alignment to flow and sprouting inhibition increase proportionally with shear stress. Scale bars: 50 μm and 100 μm . The figure has been rearranged from the original. B) Fluid simulation of the device to assess flow velocity applied to the gel with shear stresses of 5 dynes per cm^2 (top) and 15 dynes per cm^2 (bottom).¹⁷⁵ Scale bar: 200 μm . The figure has been rearranged from the original. C and D) Studies focusing on interstitial flow. C) Device setup to generate hydrostatic pressure induced by the difference in medium level.¹⁸² The $\Delta P(t)$ remains around 5 Pa, generating 10 pL h^{-1} flow corresponding to a 0.015 $\mu\text{m s}^{-1}$ velocity. D) Effect of interstitial flow and direction on angiogenesis.⁵⁷ Flow towards sprouts (S-N flow) improves sprouting (ii), while flow from the sprouts (N-S flow) inhibits angiogenesis (iii). Scale bars: 200 μm . The figure has been rearranged from the original. E and F) Compression and extension effects on vascular MPS. E) Intraluminal pressure effect on lumen morphology and cell structure.¹⁸⁹ A pressure of 100 Pa generates an inflated vessel and a change of actin morphology, into a stress fiber phenotype. Scale bars: 50 μm . F) Original lung-on-chip model with stretch modeling.²⁶ Vacuum is applied to the side chambers, generating a stretch of the central channel and attached cells. G-I) Permeability assessment in vascular MPS. G) Permeability assay in a lumen structure lined with no cells, HUVECs, or HUVECs and pericytes.¹⁹⁸ Dextran rhodamine (70 kDa) has been used over a course of 30 min. Leakages from the lumen to the surrounding ECM highly decrease when lined with both cell types. Scale bars: 500 μm . H) Permeability assay using small- to large-sized molecules in a 3D vascular network over a 12-minute course.¹⁹⁷ Permeability decreases inversely to molecule size. Scale bar: 100 μm . I) TEER method for evaluating endothelium permeability over 48 h exposure to a drug (staurosporine).²⁰⁰ Staurosporine disrupts the endothelial barrier. Scale bar: 100 μm . All figures have been edited to enhance readability.

organ-specific vascular organization has been studied, primarily in BBB models. Two studies demonstrated improved vascular bed formation and enhanced perfusability under IF conditions,^{53,184} with one also showing elevated EC marker expression and cellular alignment perpendicular to the direction of flow.¹⁸⁴ Similarly, Zhang *et al.* established two 3D self-organizing models to investigate the molecular mechanisms underlying IF-mediated vasculogenesis.¹⁸⁵ They found that matrix metalloproteinase-2 (MMP-2) was upregulated in response to IF and further explored their interactions in vasculogenesis in both HUVEC- and HBMEC-based models.

In addition to IF, another flow type is emerging in the vascular MPS field: transluminal flow.^{186,187} Offeddu *et al.* focused on the effects of transmural flow on molecular transport across the endothelial barrier, demonstrating that the flow closely mimics *in vivo*-like interstitial transport and varies according to molecular size and properties (protein *vs.* non-protein).¹⁸⁶ Meanwhile, Wang *et al.* showed that transendothelial flow not only enhanced angiogenesis of lined ECs but also promoted anastomosis with pre-existing vascular networks.¹⁸⁷ These studies are examples leading the way in the exploration of transluminal flow and its effect on vasculature processes.

Compressive/extensive forces: pressure and stretch. While shear stress and interstitial flow have been in the spotlight, other mechanical cues have also been explored using vascular MPS. Intraluminal pressure is one of the mechanical cues that is thought to play a critical role in regulating *in vivo* vascular environment, but remains poorly understood. Vascular MPS provides a promising avenue for elucidating unanswered questions about intraluminal pressure, highlighted by the study by Offeddu *et al.*,¹⁸⁶ where a 1–2 kPa intraluminal pressure generates a transmural flow and subsequent interstitial flow. Static pressure is another mechanical factor of interest, although it has been less frequently studied. Studies have shown that a static pressure of 200 Pa (1.5 mmHg) inhibits angiogenesis,¹⁸⁸ whereas a pressure of 100 Pa has been found to enhance transendothelial transport and induce vascular morphological changes¹⁸⁹ (Fig. 3E).

Additionally, the stretch of the endothelium is also of interest, as first highlighted in the landmark lung-on-chip study by Huh *et al.*²⁶ (Fig. 3F). Since that breakthrough, several vascular MPS setups using mechanical or pneumatic actuation systems have been developed, aiming to replicate and study endothelium stretching.^{190,191}

Permeability: assessment methods and applications. The permeation of molecules across blood vessel walls is crucial for the transport of nutrients and signaling molecules from the blood to surrounding tissues. Therefore, permeability has been of strong interest as an essential parameter for assessing the functionality of vasculature models, both *in vivo* and *in vitro*. Organ-specific variations in vascular permeability can be linked to endothelium types, with fenestrated vessels in the kidneys or discontinuous vessels in the liver exhibiting high permeability, whereas the BBB is characterized by lower permeability.¹⁹² Additionally, enhanced vascular permeability is often observed under inflammatory conditions,¹⁹³ underscoring its relevance in pathological contexts. These elements emphasize that permeability is a key parameter to evaluate vascular systems *in vitro*.

In the context of vascular MPS, permeability is a critical criterion for assessing the formation and functionality of vascular networks. The most widely utilized method for assessing permeability is measuring the transport of fluorescent molecules across the endothelial layer. This method is applicable for both 2D bilayer^{194,195} and 3D vessel formats.^{66,196,197} In 2D devices, permeability is typically measured by analyzing the passage of molecules from one channel to the other, offering the advantages of higher throughput and reproducibility.¹⁹⁵ In contrast, 3D devices evaluate permeability by measuring the leakage of molecules from the vascular lumen to the surrounding microenvironment. Despite being more technically challenging and presenting a lower throughput, 3D formats enable the recapitulation of more complex microenvironments, with structures involved in permeability modulation *in vitro*, such as lumen lined with pericytes¹⁹⁸ (Fig. 3G) or BBB composed of ECs in contact with astrocytes and pericytes¹⁹⁷ (Fig. 3H). Another widely used method for assessing permeability is trans-endothelial electrical

resistance (TEER). TEER measures barrier integrity by quantifying the electric current passing through the endothelial layer.¹⁹⁹ This method offers benefits such as higher time resolution, smaller experimental footprints compared to fluorescence methods, and potential scalability toward high throughput measurement^{107,200,201} (Fig. 3I).

Permeability assays have been employed to investigate the effect of chemical, physical, or biological cues on the vasculature. Cytokines and other biochemical stimuli have been used to model vascular responses under inflammatory and infectious conditions.^{38,202–205} For example, Fengler *et al.* developed an iPSC-derived microvascular model to assess changes in permeability following IL-1 β treatment, identifying anti-inflammatory compounds.³⁸ Similarly, Faley *et al.* proposed an airway-on-chip for SARS-CoV-2 studies, correlating permeability changes with viral load and other physiological readouts.²⁰⁶

Additionally, permeability in relation to other cell types or the microenvironment has also been a focus of numerous studies. In a blood-retinal barrier model *on-chip*, where retinal pigment epithelium and endothelium interact, permeability was assessed under varying degrees of epithelium injury.²⁰⁷ This study demonstrated that permeability increases with prolonged exposure to reactive oxygen species (ROS)-induced injury. Other studies focused on the impact of fibroblasts²⁰³ and pericytes^{142,198} on permeability. Among vascular MPS, the BBB remains the most extensively studied using permeability assessments. BBB models typically incorporate ECs, pericytes, and astrocytes to mimic *in vivo*-like reduced permeability,^{107,197,208} providing insights into the intricate interplay between these cell types and their role in maintaining barrier integrity.

Overall, despite their widespread use, permeability assays face several technical challenges. Fluorescence-based assays can be cytotoxic or lead to residual signals, making it difficult to follow the evolution of permeability throughout an experiment. Additionally, 3D permeability assessment mostly relies on quantification based on images, which is less direct and potentially more prone to bias than direct absorbance-based quantifications. On the other end, electrical methods such as TEER may suffer from reproducibility issues and geometric constraints, particularly when adapted to 3D formats. Emerging technologies address these limitations. For example, automated image analysis tools, such as macros, can reduce user bias in fluorescence-based assays.¹⁹⁷ Additionally, the use of electroactive tracers offers a promising hybrid approach that combines the benefits of molecular diffusion and electrical measurement.¹⁹⁴ These continued progresses for permeability assessment provide new avenues for more precise evaluations and more versatile applications.

III.2) Studies and applications for biomedical prospects

Vascular MPS offer remarkable versatility, making them a powerful tool for exploring biomedical applications. Initially

designed to mimic tissues and organs, these systems have advanced considerably over time. With the growing interest in spheroids and organoids, efforts to integrate these models into vascular MPS have expanded, enhancing their physiological relevance. Importantly, vascular MPS are not limited to replicating healthy conditions; they have also become invaluable tools for studying vascular diseases and cancer, as well as for more biomedical aspects such as immunity, therapy, and drug screening.

Tissue modeling: interface and organ-like structures. MPS based on pre-patterning methods have been developed to model various human organs, for example models of lungs,^{26,79} liver,²⁰⁹ and kidneys,^{210,211} with a focus on the interactions between endothelium and epithelium (Fig. 4A). Meanwhile, in the field of MPS based on self-organizing vascular methods, the BBB-on-chip is currently of major interest, with models integrating ECs, pericytes, and astrocytes^{197,208,212} to mimic the complex microenvironment of the BBB (Fig. 4B).

Tissue modeling has been central to MPS development since the inception of OoC technology. However, a new frontier emerges combining organoids and 3D organ modeling in MPS to create functionally vascularized human organoids. Organoids, widely used in biomedical research, recapitulate human organ structural and morphological features, as well as disease conditions *in vitro*.^{213,214} These systems are developed using classical cell lines, primary cells, or increasingly iPSCs.²¹⁵ Despite their ability to accurately mimic human organ structure, organoids face critical limitations due to the absence of well-defined vascular networks.^{215–217} Without perfusable vasculature, nutrient and oxygen delivery to the inner regions of organoids is restricted, resulting in size limitations and necrotic cores. Furthermore, the lack of functional perfusable vasculature diminishes their potential use as drug screening platforms. *In vivo* transplantation of organoids has successfully induced vascularization,^{218,219} however, efforts are now focused on overcoming these limitations *in vitro* by leveraging MPS technologies to vascularize organoids.

When discussing vascularization in MPS, we will classify organoids into two categories: spheroids and proper organoids. Spheroids are aggregates of either healthy or tumor-derived cells – usually cell lines or primary cells – aggregated in a disorganized structure, which can be mixed with other cell types (ECs, fibroblasts, *etc.*). Proper organoids are composed of healthy cells derived from pluripotent stem cells and exhibit organized organ-like structures. Spheroids are generally easier to vascularize by mixing target cells and ECs, aiming to promote anastomosis with pre-existing vascular networks or angiogenic sprouts. For instance, Jin *et al.* generated hepatic spheroids by co-culturing induced hepatic cells with HUVECs, achieving an even higher degree of vascular development when exposed to flow²²⁰ (Fig. 4C). Similarly, Bonanini *et al.* successfully obtained two types of vascularized hepatic spheroids *on-chip*, with vasculature

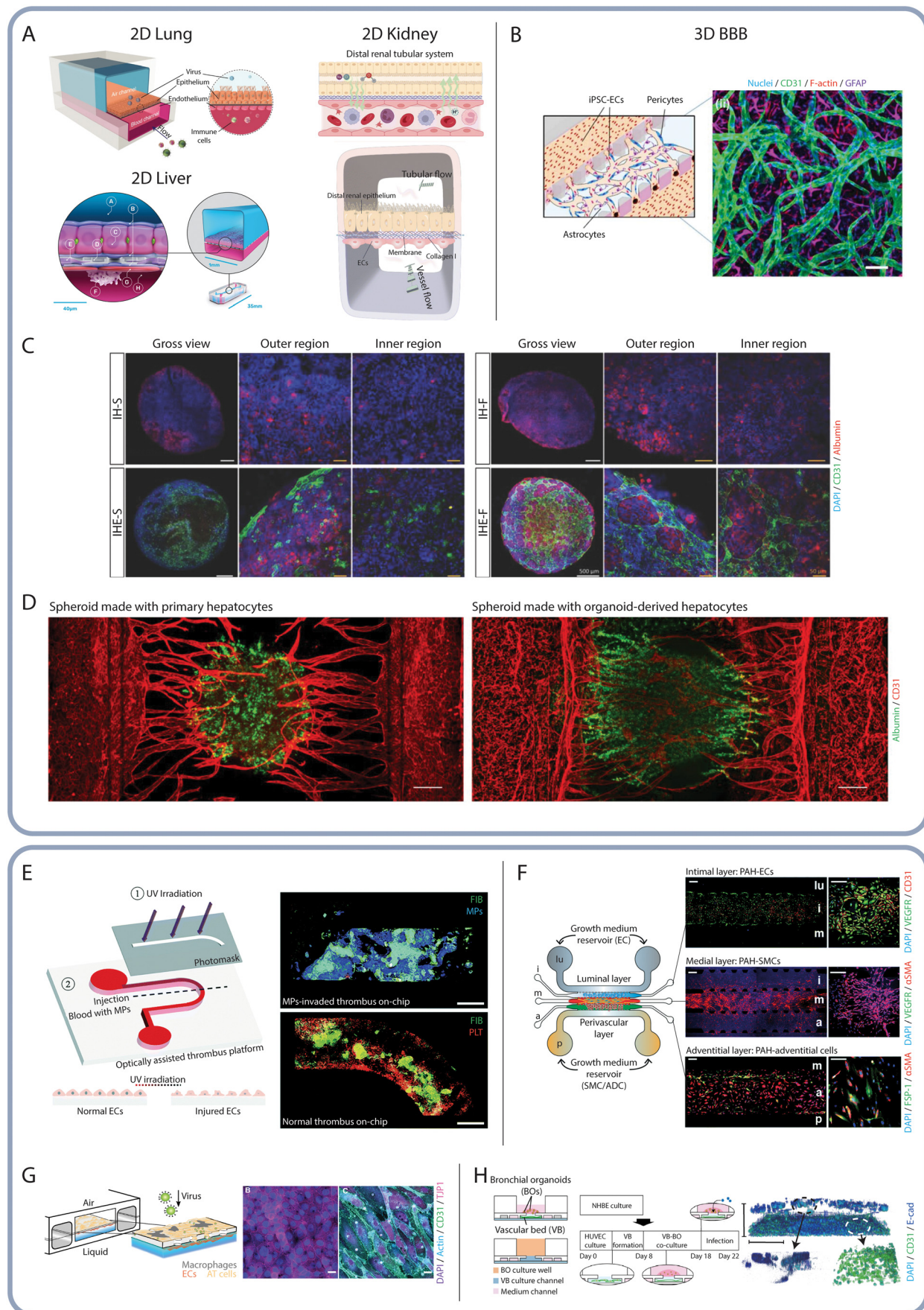


Fig. 4 Vascular MPS for tissue modeling and organ development. A) Schematics of various 2D bilayer models such as lung-, liver- and kidney-on-chip co-culturing organ-specific epithelial tissue and ECs.^{79,209,211} B) Vascular bed *on chip* schematic and confocal image. hiPSC-derived ECs, brain pericytes, and astrocytes formed a vascular network (ii), recapitulating the *in vivo* 3D interface of the human BBB.²⁰⁸ Scale bar: 100 μ m. C) Induced hepatic (iHep) cells were cultured with or without HUVECs (iHE or iH), under static or flow conditions (iHE-S/iH-S or iHE-F/iH-F), producing a vascularized organoid-like tissue.²²⁰ White scale bars: 500 μ m, yellow scale bars: 50 μ m. D) Vascularized hepatic spheroids made of primary hepatocytes and HUVECs (left) and fetal primary hepatocytes from hepatic organoids (right), using an angiogenesis method.¹⁴⁰ Scale bars: 200 μ m. E) Thrombosis-on-chip model mimicking injury-induced thrombosis, using whole blood and microparticles.²²⁸ Scale bars: 50 μ m. F) PAH-on-chip encapsulating pulmonary arterial ECs, SMCs, and adventitial cells, organized in a three-layer configuration.²³⁵ Scale bars: 250 μ m and 100 μ m. The figure has been rearranged from the original. G) 2D bilayer device recapitulating endotheliitis and vascular damage after SARS-CoV-2 infection.⁷⁸ B and C are the epithelial and endothelial layers, respectively. Scale bars: 20 μ m. H) Vascular network-bronchial organoid *on-chip* used to mimic vascular damage induced by infected lung epithelium paracrine signaling.⁵⁶ Scale bars: 500 μ m. The figure has been rearranged from the original. All figures have been edited to enhance readability.

infiltrating the spheroids¹⁴⁰ (Fig. 4D). One method involved combining primary hepatocytes with HUVECs, while the other consisted of generating hepatic organoids from fetal primary cells, dissociating them, and reaggregating the cells into spheroids without EC mixing. Other studies have also demonstrated vascularization in cortical²²¹ or liver²²² spheroids, even in the absence of direct combination with ECs.

In contrast, vascularizing proper organoids *in vitro* remains more challenging. Achieving functional vascular integration often requires extensive optimization, including synchronization of co-culture timing, selection of appropriate supporting cell types, and modulation of biochemical and physical cues.²¹⁷ For many organ systems, generating fully perfusable and functional vasculature within proper organoids is still an unmet goal.²¹⁶ However, recent progress has identified potential molecular targets to support vascular development. For example, Shaji *et al.* identified CYR61 and HDGF as promising regulators of vascularization in cerebral organoids, offering new strategies to improve vascularization *in vitro* perfusion and maturation.²⁰⁵ Together, these efforts highlight the increasing potential of MPS platforms to overcome vascularization challenges in both spheroid and proper organoid models, moving the field closer to fully functional, physiologically relevant tissue constructs.

Vascular diseases: specific and non-specific to the vasculature. MPS have proven to be powerful tools for studying vascular diseases, affecting either an organ, blood, the vasculature itself, or a combination.²²³ Pathologies related to blood and vasculature, such as thrombosis and hypertension, have been explored using vascular MPS. Moreover, despite the existence of vascular-specific diseases, the endothelium can also be altered by pathologies specific to neighboring tissues. The example of viral infections has successfully highlighted the effect of infections on vasculature, using organ-specific vascular MPS.

Thrombosis is the aggregation of platelets, red blood cells, and fibrin within blood vessels and poses a significant risk for heart attacks and strokes if unresolved. Thrombosis-on-chip models have been developed for disease modeling and anti-coagulant drug screening.^{40,224-228} For example, Barrile *et al.* demonstrated thrombus formation triggered by an anti-CD154 monoclonal antibody,²²⁶ while Chen *et al.* showed the induction of thrombosis due to microplastic exposure in a

vascular MPS model²²⁸ (Fig. 4E). Outside of PDMS soft lithography MPS, advancements in 3D printing and bioprinting technologies have had a huge impact for thrombosis-on-chip modeling, enabling a more accurate vessel architecture in devices.^{229,230}

Hypertension is characterized by elevated blood pressure, defined clinically as a systolic blood pressure (SBP) higher than 130 mmHg and a diastolic blood pressure (DBP) exceeding 80 mmHg.²³¹ Histologically, hypertension is associated with vascular remodeling, which typically manifests as inward and hypertrophic changes, involving a reduction in luminal diameter due to SMC hyperplasia and hypertrophy within the tunica media.^{232,233} Despite the global burden of hypertension, vascular MPS models remain relatively limited, with a few examples focused on PAH. PAH specifically affects pulmonary arteries, defined by a mean pulmonary arterial pressure (mPAP) above 20 mmHg, and is similarly characterized by vascular remodeling.²³⁴ Two studies have successfully recapitulated PAH-related wall thickening and muscularization using MPS. Al-Hilal *et al.* developed a three-layer PAH model incorporating ECs, SMCs, and adventitial cells to mimic the tunica intima, media, and externa, respectively²³⁵ (Fig. 4F). They applied flow-induced mechanical stress and succeeded in reproducing vascular remodeling features such as wall thickening and muscularization. In parallel, Ainscough *et al.* used a two-layer PAH model, using ECs and bone morphogenetic protein receptor type II (BMPR2)-mutated SMCs, combined with hypoxia.³¹

Beyond hypertension and thrombosis, other vascular pathologies have also been explored using MPS, including arteriovenous malformation (AVM)-on-chip²³⁶ and atherosclerosis-on-chip,²³⁷ further emphasizing the potential of vascular MPS to study both vascular-specific and vascular-involved diseases in controlled environments. Additionally, while 2D bilayer devices are widely used to emulate viral infections targeting epithelial cells,²³⁸ recent studies have shown that viral infections can affect the vasculature in addition to their primary organ targets, notably SARS-CoV-2.²³⁹ In 2D device studies, a few models successfully highlight the immune response activation in ECs mediated by signals from SARS-CoV-2-infected epithelial cells^{78,240} (Fig. 4G). A similar study has also revealed region-specific EC immune responses in influenza virus-infected

respiratory tracts.²⁹ Furthermore, a few 3D models have replicated similar immune and vascular responses after SARS-CoV-2 infection, including lung-on-chip⁵⁶ (Fig. 4H) and heart-on-chip²⁴¹ models.

Cancer: tumor vasculature and metastasis. Tumor-induced angiogenesis and vascularization are major hallmarks of cancer development.^{5,6} To achieve angiogenesis, cancer cell lines can be used to generate growth factor gradients and enhance sprouting. For example, the glioblastoma cell line (U87MG) has been shown to produce VEGF, effectively inducing angiogenesis.²⁴² Once established, tumor vasculature differs markedly from healthy vascular networks, exhibiting increased heterogeneity, abnormal permeability, multidirectional flow, and disorganized distribution.²⁴³ Accurately modeling tumor vasculature is therefore key for developing functional *in vitro* tumor models.²⁴⁴

The vascularization of tumor spheroids has recently gained momentum in the MPS field.^{69,245-247} Nashimoto *et al.* engineered a perfusable and vascularized tumor spheroid by co-culturing HUVECs, breast cancer cells (MCF-7 line), and human lung fibroblasts.⁶⁹ Vascularization was achieved by spheroid-induced angiogenesis directed toward the spheroid, followed by anastomosis, as confirmed by immunostaining and histological analysis. Additionally, the anti-cancer effects of paclitaxel on both the spheroid and vasculature were assessed. Two other studies focused on alveolar soft part sarcoma (ASPS) also achieved intra-spheroid vascularization by inducing angiogenesis directed toward the spheroid^{245,246} (Fig. 5A). Alternatively, vasculogenesis methods can be used to generate surrounding vasculature, which would then penetrate the spheroids. Straehla *et al.* generated a vascularized glioblastoma by co-culturing a tumor spheroid and a BBB vascular network, exploring permeability and drug therapy considerations.²⁴⁷ In contrast, Haase *et al.* developed a model in which tumor spheroids were encapsulated within a vascular MPS, focusing not on intra-spheroid vascularization but on the tumor-specific vasculature signature.²⁴⁸ This approach emphasized the tumor microenvironment and its drug-resistance properties. It is noteworthy that vascularization was observed after extended culture times, but was not the primary objective of the study.

Metastasis is another hallmark of cancer, which consists of the sequential processes of cancer cell intravasation from the primary tumor into the vessels, migration through the vasculature, and extravasation into the secondary sites.^{5,6} Tumor intravasation is influenced by multiple factors, including chemotactic gradients, oxygen levels, and impaired endothelial barriers.²⁴⁹ For example, the transmigration of aggressive breast cancer cells (MDA-MB-231 cell line) across the endothelial layer has been observed under normal conditions, and was further enhanced after TNF- α treatment.²⁴² Similarly, Zervantonakis *et al.* explored the role of macrophages in intravasation, finding that macrophage-secreted TNF- α increases endothelial permeability, thereby enhancing the intravasation of fibrosarcoma cells²⁵⁰ (Fig. 5B). Extravasation has been widely studied in both

2D^{251,252} and 3D formats.²⁵³ Jeon *et al.* modeled the organ-specific metastasis of breast cancer cells into a bone-like environment.²⁵³ Several studies have focused on identifying parameters that facilitate extravasation. For example, a study shows that tumor cells physically trapped within blood vessels exhibit higher trans-endothelial migration efficiency compared to non-trapped cells, with similar findings for cell clusters *versus* individual tumor cells²⁵⁴ (Fig. 5C). Another study demonstrated that hypoxic conditions increase extravasation, as shown using three different breast cancer cell lines.²⁵⁵

However, as both intravasation and extravasation predominantly occur in capillary-scale vessels, physiologically relevant vascular networks are required to obtain improved metastasis models. To accurately mimic both the tumor site and metastatic niche, vascular MPS should reflect not only capillary dimensions but also organ-specific endothelial characteristics. Indeed, Miles *et al.* discussed two complementary mechanisms of extravasation: the trapping model, where cells become physically trapped in capillaries, and the adhesion model, where tumor cells bind to specific endothelial markers in larger vessels.²⁵⁶⁻²⁵⁸ Their findings underscore the critical roles of both vessel diameter and endothelial phenotype in regulating tumor cell arrest and transmigration. While a HUVEC-based model has successfully recapitulated tumor cell trapping *on-chip*,²⁵⁴ 3D organ-specific microvascular models are required to investigate the relative contributions of adhesion *versus* trapping across different tumor types and metastatic niches. This limitation further highlights the previously discussed gap in self-organizing vascular MPS using organ-specific microvascular ECs and underscores the need for future studies to develop and apply such models for metastasis research.

Immunity: immune responses and therapy. Vascular MPS has also been used to model immune responses, as blood vessels are not only responsible for oxygen or nutrient delivery but also for transporting circulating cells such as platelets and immune cells. In the context of immunity, platelets are primarily associated with coagulation following vascular injury. While many studies have focused on platelet aggregation in the setting of thrombosis, hemostasis-on-chip models have also been developed to investigate bleeding responses.²⁵⁹ These models replicate bleeding events using different methods, such as valve actuation to mimic a wound opening²⁶⁰ or by needle puncture to mimic injuries,²⁶¹ both of which induce hemostasis. Additionally, the role of macrophages in coagulation has been explored using vascular MPS, further highlighting the complex interplay between immune cells and hemostatic processes.³⁶

Beyond platelets, immune cells such as neutrophils, monocytes, dendritic cells, NK cells, and T cells play critical roles in immune surveillance and pathogen elimination. Neutrophils, as part of the innate immune system, serve as the first line of defense by targeting circulating pathogens within the vasculature through mechanisms such as

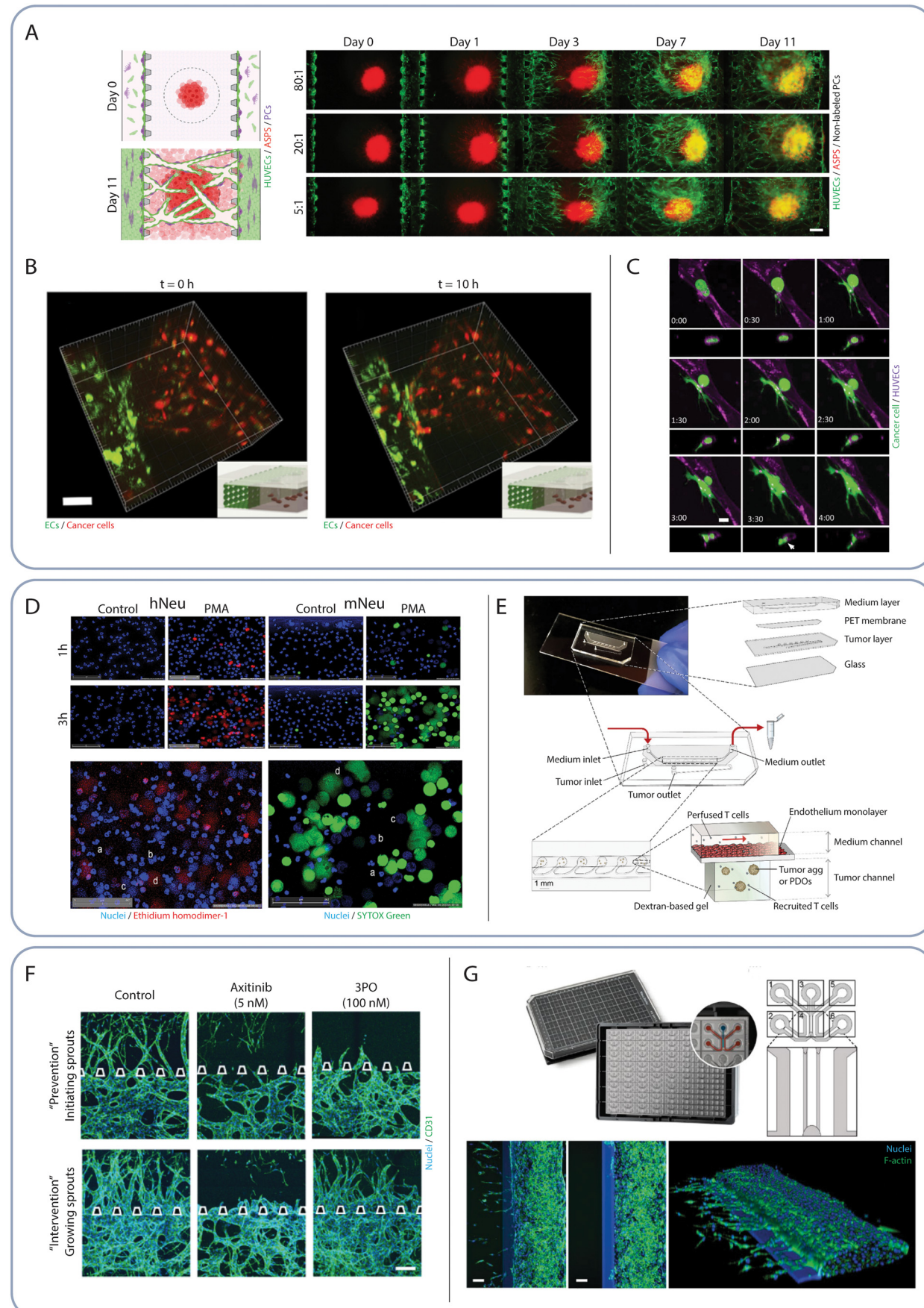


Fig. 5 Vascular MPS for cancer recapitulation, immunity modeling, and drug screening. A) Alveolar soft part sarcoma (ASPS) model *on-chip*, using an angiogenesis method to vascularize the tumor spheroid.²⁴⁶ Scale bar: 200 μm . B) Intravasation of fibrosarcoma tumor cells (HT1080, red) from the ECM through the HUVEC monolayer (green) at 0 h and 10 h.²⁵⁰ Inset shows image orientation in the device. Scale bar: 50 μm . C) Extravasation of a tumor cell (MDA-MB-231, green) from the endothelium (purple).²⁵⁴ Scale bar: 10 μm . D) NET formation *on-chip* from human (hNeu) and monkey (mNeu) neutrophils, activated by 100 nM phorbol 12-myristate 13-acetate (PMA). Green is SYTOX-green, and red is ethidium homodimer-1, revealing DNA's presence. Letters a to d highlight different stages of NET formation.²⁶² The figure has been rearranged from the original. E) Device setup for immunotherapy *on-chip* using CAR-T cells targeting breast tumor cells, contained either in tumor aggregate or patient-derived organoids.²⁷¹ F) Drug assay, using 5 nM axitinib or 100 μM 3PO, against angiogenic sprouts.⁵⁷ Either prevention or intervention methods have been applied, where drugs are injected before or during sprout formation, respectively. Scale bar: 200 μm . The figure has been rearranged from the original. G) A high-throughput device used to perform angiogenesis phenotypic screening in response to various types of drugs.²⁸¹ Scale bars: 100 μm . The figure has been rearranged from the original. All figures have been edited to enhance readability.

neutrophil extracellular traps (NETs). Yadav *et al.* demonstrated the utility of MPS for studying NET formation using pig ECs and blood²⁶² (Fig. 5D). Once pathogens are detected in tissues, immune cells extravasate from blood vessels into infected regions. MPS have replicated these processes, including monocyte extravasation,²⁶³ dendritic cell migration,²⁶⁴ and natural killer (NK) cell activity.^{265,266} Ayuso *et al.* modeled NK cell targeting of solid tumors, supported by circulating antibodies,²⁶⁵ while Humayun *et al.* targeted *Toxoplasma gondii*-infected cells.²⁶⁶ Similarly, T cells' adhesion, crawling, and extravasation have been observed in a 3D vascular network.⁷⁰

Immunity-on-chip has become of major interest in the MPS field, with efforts focused on developing both lymph node-on-chip models and integrated *on-chip* vascular-lymphatic networks. Analogous to angiogenesis, the role of flow has been demonstrated to enhance lymphatic vessel sprouting.²⁶⁷ Additionally, lymphatic endothelial tubules have been used to explore the effect of shear stress, further advancing our understanding of lymphatic biology.²⁶⁸ While standalone lymphatic vessel and lymph node *on-chip* models are still under development, the intrinsic relationship between blood and lymphatic vessels suggests that next-generation MPS will likely combine these two networks.^{269,270} Such integrated systems could provide more comprehensive investigations of immune dynamics, including cell trafficking, antigen presentation, and systemic immune regulation.

From a biomedical and translational perspective, vascular MPS are increasingly employed in immunotherapy research, particularly in the evaluation of chimeric antigen receptor (CAR)-T cell therapies. For instance, Maulana *et al.* reported an *on-chip* breast cancer model to assess the efficacy of CAR-T cells in endothelial transmigration, tumor infiltration, and immune activation²⁷¹ (Fig. 5E). Similarly, CAR-T cells have been used in a vascularized *on-chip* model of ovarian tumor spheroids, derived from the Skov3 cell line.²⁷² This study shows that the spatial organization of the spheroid and, ultimately, the tumor architecture significantly impact therapeutic efficiency. Spheroids surrounded by fibroblasts exhibited higher vascularization and perfusability, compared to those with fibroblasts embedded within, which ultimately allowed an enhanced CAR-T cell delivery and more effective therapeutic assessment. Other models have also been designed to study CAR T-cell responses in ovarian and liver cancer models.^{273,274}

Drug screening: low and high throughput. Given the critical role of vasculature in delivering drugs to target tissues, vascular MPS emerged as a powerful tool for drug-related studies. Additionally, the enactment of the Food and Drug Administration (FDA) Modernization Act 2.0 has further encouraged the use of non-animal models, opening new opportunities for MPS in drug testing and screening.²⁷⁵ When considering *in vitro* platforms for drug evaluation, two parameters must be balanced: complexity and throughput. Complexity refers to how closely a model replicates the structure and function of real human tissues, while throughput refers to the number of samples that can be processed in a single experimental run. These parameters are often inversely related: highly complex models typically support low-throughput, whereas high-throughput platforms require simplification to maintain scalability. The appropriate balance depends on the research objective: low-throughput, high-complexity models are commonly used in academic settings for mechanistic studies, while high-throughput, lower-complexity models are favored in industry for efficient drug screening.

Low-throughput vascular MPS, while more intricate, are ideally suited for detailed analyses of drug delivery, efficacy, and mechanisms of action. These models typically assess one or a few compounds at a time, offering fine-grained insights into how drugs interact with tissue barriers and vasculature. For example, Kim *et al.* demonstrated that vascularizing lung tumor spheroid-on-chip enhanced both drug delivery and immune cell infiltration.²⁷⁶ Using 2D devices, some studies assessed the transport of nanocarriers across endothelial barriers, such as the BBB.⁵¹

Drug assays in low-throughput vascular MPS usually aim to disrupt vascular networks, especially in cancer research, to target tumor vasculature. Studies often focus on disrupting already-formed vessels^{48,277} or inhibiting angiogenesis.⁵⁷ Kim *et al.* tested two anti-angiogenic drugs, axitinib and 3PO, showing that axitinib inhibits both initiation and elongation of vascular sprouts, while 3PO is effective only during the initiation phase⁵⁷ (Fig. 5F). Similarly, vascularized organoid-on-chip models have been employed in drug assays, although there is still room for improvement.²⁷⁸ One such study developed colorectal and breast cancer tumor spheroids, with both surrounding and penetrating vascular structures, to evaluate various drugs such as laninifab and cabozantinib. This study highlights their effectiveness in disrupting pre-

existing vasculature.⁴⁸ In contrast, some drug assays aim to enhance vasculature or restore functionality when vascular dysfunctions occur. For example, brain arteriovenous malformation, causing impaired vascular functionality, has been modeled using a vascular MPS and used for drug screening purposes.²³⁶

While traditional vascular MPS offer high functionality and relevance for drug testing, their inherently low throughput poses a major barrier to large-scale pharmaceutical screening. Addressing these challenges will be crucial for the broader adoption of MPS in industry. Towards that aim, multiplex systems have been developed to enable high-throughput microfluidic devices for vascular applications. These systems aim to balance physiological fidelity with scalability.^{37,279} A first possible application is to perform vascular phenotypic screening, focusing on parameters such as angiogenic sprout morphology²⁸⁰ and angiogenesis dynamics²⁸¹ (Fig. 5G). Secondly, disease-specific drug screening can be performed to determine potential treatments. Phan *et al.* developed a platform consisting of multiple vascularized micro-organs in a 96-well format, allowing for blind screening of a compound library, including anti-cancer drugs.²⁷⁹ Wevers *et al.* demonstrated the application of a 384-well high-throughput BBB model, integrating 40 or 96 microfluidic chips for parallel screening.³⁷ Similar approaches have also been applied to study drug responses in fibrotic conditions²⁸² and aortic aneurysm.²⁸³

Conclusion

Vascular MPS encompass diverse designs, ranging from 2D bilayer devices replicating endothelial–epithelial interfaces to 3D pre-patterned devices reconstructing vascular architecture, and self-organizing platforms which form *in vivo*-like networks through biological processes such as vasculogenesis and angiogenesis. While PDMS-based vascular MPS have proven highly effective in recapitulating vascular architecture, some limitations remain, particularly in modeling the wide range of vessel size and geometry. At the micro-scale, fabrication constraints restrict pre-patterned models, whereas self-organizing networks often yield vessels with diameters significantly larger than native capillaries. Typically less than 10 µm *in vivo*, *in vitro* vessels commonly range from 20 to 80 µm or more, especially in perfusable networks. As discussed previously, organ-specific microvascular ECs represent a promising approach to achieving capillary-scale modeling under both physiological and pathological conditions. For larger-scale vessels, PDMS-based fabrication relying on soft lithography is limited to rectangular, micrometer-scaled, and linear vessels. In contrast, native blood vessels range from micrometric to millimetric dimensions, with circular cross-sections and complex topologies. To address these limitations, 3D printing technologies have demonstrated significant potential in fabricating anatomically relevant vascular MPS.^{20–25}

Blood vessels are complex structures composed of multiple cell types and are surrounded by a specific microenvironment. Capturing such complexity remains a common challenge for *in vitro* models. However, vascular MPS are steadily advancing in biological relevance. The first major improvement lies in cell sourcing. Vascular MPS now utilize more relevant cell types, such as organ-specific ECs or progenitor cells, shifting away from the conventional HUVECs toward more *in vivo*-like endothelium. The second advancement is the incorporation of perivascular cells, such as fibroblasts, pericytes, and SMCs. These additions enhance the stability and functionality of vascular MPS, both in healthy and diseased models. Recent years have also seen a growing interest in including other cell types, such as astrocytes, neurons, immune cells, and blood cells. The third improvement involves the use of *in vivo*-like acellular components. Membrane designs have evolved towards thinner membranes approximating basement membrane dimensions and electrospun membranes mimicking *in vivo* fibrillar organization. Additionally, ECM materials have transitioned from traditional hydrogels to more complex components, including dECM and synthetic polymers. Overall, as new sources and technologies for both cellular and acellular components continue to emerge, vascular MPS are poised to achieve greater biological relevance.

Current vascular MPS have demonstrated substantial capabilities for studying a broad range of vascular biology. Physical factors play a critical role in vascular biology, influencing blood vessel formation, stability, and functionality. For instance, intraluminal flow and resulting shear stress are key regulators of vascular organization and integrity, while interstitial flow has been strongly linked to the regulation of angiogenesis. Other mechanical factors, such as pressure and stretch, have also been recapitulated in vascular MPS, though they remain relatively understudied. An essential functionality of the endothelium is permeability, which varies depending on the organ and EC type.

Despite the current implementation of physical cues in vascular MPS, significant limitations persist for two main reasons: the lack of comprehensive theoretical values and the difficulty in experimental setups. Knowledge of *in vivo* values and physiological phenomena is still limited. For example, shear stress or permeability values are often derived from *in silico* simulations rather than empirical measurements, and data on interstitial flow, intraluminal pressure, and stretch are scarce in the literature. To address these gaps, theoretical advancements are necessary and should be thereafter integrated into vascular MPS studies. Simultaneously, experimental setups required to simulate these physical cues are often complex or labor-intensive. Developing more user-friendly or ready-to-use systems will be critical to enabling the routine incorporation of such physical factors into vascular MPS models.

In addition to studies on physical parameters, vascular MPS have enabled extensive exploration of biological aspects of vascular systems. These models have evolved from 2D co-

culture systems to more advanced 3D platforms, integrating vasculature with spheroids and organoids. In the context of tumor models, a vascular bed surrounding the spheroid is already a critical tool for studying the tumor microenvironment, vascular remodeling, and phenotypic changes in the endothelium. Yet, the need for intra-spheroid vascularization remains critical, as tumoral vasculature often presents distinct biological profiles and functional behaviors,²⁸⁴ directly influencing drug delivery and therapeutic efficiency.²⁸⁴ In organoids, the lack of perfusable vasculature leads to the formation of necrotic cores, as diffusion alone is insufficient to deliver nutrients to the center. Intra-organoid vascularization is thus essential for supporting further growth, preventing necrosis, and enabling the formation of more complex and mature structures. Overall, while integrating vasculature within organoids *on-chip* remains a significant challenge, progress in the next-generation organoids and vascular MPS is expected to make this integration increasingly feasible.

Finally, beyond physiological and organ-specific modeling, vascular MPS provide a robust platform for studying pathological conditions and developing clinical applications. These models have successfully modeled a wide range of vascular disorders *in vitro*, including thrombosis, hypertension, atherosclerosis, genetic vascular diseases, and infections. Their role in biomedical applications has further expanded to include immune response studies and drug testing. This translational relevance has been validated by the U.S. FDA's recognition of MPS for use in clinical development under the FDA Modernization Act 2.0, which allows MPS to serve as an alternative to animal models in defined contexts.²⁷⁵ A remaining limitation, however, is throughput, particularly for PDMS-based MPS, which, despite offering higher throughput than animal models, still lag behind compared to 2D dish culture models. Encouragingly, recent developments in both academic and commercial MPS platforms, incorporating PDMS and alternative materials, have demonstrated promising improvements in throughput. While these innovations open doors for both vascular research and high-content drug screening, ongoing improvements in scalability and integration need to be sustained to fully establish vascular MPS as a standard tool in clinical and pharmaceutical research.

Overall, vascular MPS are highly valuable tools for studying a wide range of vascular-related physiological and pathological processes. Despite their current limitations, ongoing advancement in fabrication techniques, cell sourcing, acellular environment engineering, and technological innovation are poised to establish vascular MPS as a gold standard for vascular research.

Data availability

No primary research results, software or code have been included and no new data were generated or analyzed as part of this review.

Author contributions

Conceptualization: L. B., A. K., Y. S., H. Z., K. F., and R. Y. conceptualized the study. Funding acquisition: K. F. and R. Y. acquired the funding. Project administration: L. B. and R. Y. managed the project. Supervision: L. B. supervised the project. Visualization: L. B. designed the figures. Writing – original draft: L. B., A. K., Y. S., H. Z., and K. F. wrote the original draft. Writing – review & editing: L. B., A. K., Y. S., H. Z., K. F., and R. Y. reviewed and edited the manuscript.

Conflicts of interest

All authors have agreed upon the publication of this manuscript. R. Y. is one of the founders and shareholders of Physios Biotech, Inc.

Acknowledgements

This study was partially supported by the Japan Agency for Medical Research and Development: AMED-MPS (JP22be1004204 and JP17be0304205), AMEDP-PROMOTE (JP22ama221206), AMED P-CREATE (JP20cm0106277), and AMED-CREST (21gm1610005). The study was also supported by the Japan Society for the Promotion of Science: KAKENHI (JP24H00404, Grant-in-Aid for Scientific Research (A)), and KAKENHI (24K23941, International Leading Research: Creating a kidney). Finally, the study was supported by the "Advanced Research Infrastructure for Materials and Nanotechnology in Japan (ARIM)" of the Ministry of Education, Culture, Sports, Science and Technology (MEXT, JPMX1222KT1172). Illustrations were created with <https://Biorender.com>.

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