Abstract | Vascular endothelial cells form the inner layer of blood vessels where they have a key role in the development and maintenance of the functional circulatory system and provide paracrine support to surrounding non-vascular cells. Technical advances in the past 5 years in single-cell genomics and in vivo genetic labelling have facilitated greater insights into endothelial cell development, plasticity and heterogeneity. These advances have also contributed to a new understanding of the timing of endothelial cell subtype differentiation and its relationship to the cell cycle. Identification of novel tissue-specific gene expression patterns in endothelial cells has led to the discovery of crucial signalling pathways and new interactions with other cell types that have key roles in both tissue maintenance and disease pathology. In this Review, we describe the latest findings in vascular endothelial cell development and diversity, which are often supported by large-scale, single-cell studies, and discuss the implications of these findings for vascular medicine. In addition, we highlight how techniques such as single-cell multimodal omics, which have become increasingly sophisticated over the past 2 years, are being utilized to study normal vascular physiology as well as functional perturbations in disease.

The development of functional vasculature is crucial for the health of every cell in the body. Endothelial cells, which line the inside of all blood and lymphatic vessels, have key roles in delivering oxygen and nutrients, regulating blood flow, modulating immune cell trafficking and maintaining tissue homeostasis. Vascular endothelial cell dysfunction is central to the progression of most chronic conditions, as well as ischaemic heart disease and stroke, the top two global causes of mortality. However, many questions remain about the fundamental biology of endothelial cells; for example, how do endothelial cell progenitors differentiate into distinct subtypes; how do they achieve organotypic heterogeneity; and how is the function of specialized endothelial cells perturbed in disease? In the past 5 years, the advent of single-cell RNA sequencing (scRNA-seq) technologies, combined with increasingly sophisticated strategies for genetic lineage tracing, have allowed the field to address these questions. In this Review, we highlight new discoveries that can be generalized to blood vessel development across all tissues, such as the role of cell cycling in the differentiation of endothelial cell subtypes as well as the phenomenon of ‘pre-artery’ specification. In addition, we discuss new findings for organ-specific endothelial cells in several key organs such as the heart, brain, lungs, kidneys and liver. Lymphatic endothelial cell development and heterogeneity have been reviewed elsewhere and are beyond the scope of this Review. While this Review is not intended to comprehensively cover all aspects of organ-specific vasculature, we aim to demonstrate the wealth of new insights into endothelial cell biology that can be gained from scRNA-seq and integrated multimodal omic atlases. Particularly groundbreaking discoveries include the identification of functionally distinct capillary subpopulations in the lung, kidney and liver, as well as capillary zonation in the brain and heart. The identification of novel endothelial cell subtypes and molecular markers, as well as cell–cell interactions, has implications for tissue engineering and the development of novel therapeutic strategies for cardiovascular disease.

Arterial and venous differentiation

During embryogenesis, blood vessels must first arise de novo in a process known as vasculogenesis. The differentiation of angioblasts (endothelial cell progenitors) and their transition to endothelial cells are tightly regulated by transcription factors and signalling pathways that have been extensively studied. The first blood vessels arise from the rapid proliferation of endothelial cells, which subsequently form vascular tubes to create separate arteries and veins. Much research over the past two decades has described the crucial role of vascular endothelial growth factor (VEGF) and Notch signalling during arteriovenous differentiation. In the broader context, increased VEGF and Notch signalling...
**Key points**

- Arterial differentiation is coupled to cell cycle arrest, and new arteries expand during development by recruiting endothelial cells from capillaries and veins.
- Single-cell RNA sequencing (scRNA-seq) data and multimodal omics atlases provide insights into the heterogeneity of endothelial cells across tissues, as well as conserved transcription codes during development.
- Novel methods of endothelial cell lineage tracing combined with scRNA-seq has facilitated the identification of functionally distinct capillary populations in the lung, liver and kidney, as well as their transcriptional response to disease.
- Capillary zonation, or the gradual phenotypic continuum of endothelial cells along an axis, has been identified in multiple organs including the brain, heart and liver, and communication between endothelial cells and mural cells has a key role in maintaining capillary zonation and important structures such as the blood–brain barrier.
- Comparison of scRNA-seq atlases from humans and mice reveals that endothelial cell populations are largely conserved between species across multiple organs and that although unique species-specific gene expression patterns exist for endothelial subtypes, the high degree of similarity between species demonstrates the utility of mice as a model organism for vascular biology.
- Advances in bioengineering have led to the creation of organoids with increasingly functional vasculature; key insights from single-cell studies are expected to facilitate improved differentiation protocols for human pluripotent stem cells as well as the in vitro development of organ-specific vasculature.

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**Reviews**

Single-cell RNA sequencing (scRNA-seq) data and multimodal omics atlases provide insights into the heterogeneity of endothelial cells across tissues, as well as conserved transcription codes during development.

**Cell cycling and endothelial cell fate**

Insights from scRNA-seq and mosaic analyses have spurred the advances in our understanding of how Notch and COUP-TFII signalling support and restrict arterial differentiation, respectively. Surprisingly, the mechanism seems to involve the effects of Notch and COUP-TFII signalling on cell cycling and metabolism, rather than the direct regulation of arterial and venous cell fate determinants. The first study describing the relationship between cell cycling and arterial differentiation used the retina as a model system. During blood vessel formation, initiation of blood flow resulted in shear stress-induced Notch signalling and subsequent upregulation of gap junction α4 protein (also known as connexin 37). Gap junction α4 protein signalling led to flow-induced endothelial cell cycle arrest via p27 expression, enabling the expression of arterial specification genes. A similar relationship between arterial differentiation and cell cycling was subsequently described in the developing heart, although in the heart, arterial differentiation was initiated in single endothelial cells within the immature vascular plexus before the initiation of blood flow. These individual pre-artery endothelial cells were lineage-traced to mature coronary arteries, establishing these endothelial cells as pre-specified progenitors. Pre-arterial differentiation was augmented by Notch activation, but blocked by COUP-TFII overexpression, owing to the transcriptional activation of cell cycle genes.

Numerous epigenetic studies support the link between cell cycle genes and the inhibitory effects of COUP-TFII on arterial differentiation. In a genome-wide analysis comparing COUP-TFII DNA binding with artery-specific and vein-specific transcriptional enhancers, COUP-TFII was over-represented at vein-specific enhancers, the majority of which were near actively expressed cell cycle genes. The association between proliferating endothelial cells and capillary clusters with enriched expression of genes characteristic of venous endothelial cells had previously been identified in the brain as well as in regenerating aortas, adding further evidence supporting the idea that actively cycling cells are skewed towards a venous fate. Investigators are also beginning to identify the molecular mechanisms underlying the role of the cell cycle in arterial endothelial cell differentiation. Genomic analyses of mice expressing the fluorescent ubiquitination cell cycle indicator reporter specifically in endothelial cells showed that endothelial cells in the late G1 state are more sensitive to transforming growth factor-β1 (TGFβ1)-stimulated arterial specification-associated gene transcription than those in the early G1 state. These new studies correlating arterial endothelial differentiation with cell cycle exit contribute to our mechanistic understanding of arterial differentiation during development and form an important embryological parallel to older observations that mature arterial endothelial cells rarely proliferate in adults.

Pre-specification of arterial endothelial cells has also been described in other vascular beds such as the zebrafish trunk and the mouse developing retina in cell populations with low proliferative rates and high Notch activity. Live imaging in zebrafish revealed activation of Notch in tip cells, which upregulated cxcr4a (encoding the chemokine C-X-C motif receptor 4a) expression in endothelial cells, enabling them to connect with and join arterial vessels. In the mouse retina, a subset of cells within the tip cell subpopulation similarly receives higher Notch signalling, which facilitates arterial differentiation (as indicated by the expression of Cxcr4) and migration of cells to form arteries. Pre-specification at the tip cell location is supported by data from scRNA-seq analyses showing that tip cells and arterial endothelial cells are closely related at the transcriptional level. In addition to Notch, increased VEGF–ERK signalling in both the retinal tip cells and coronary vessel plexus is another upstream trigger of cell cycle exit and arterial differentiation. VEGF is normally expressed where arterial pre-specification occurs, notably the retinal tip cell region and the intramyocardium. A mosaic analysis that involved genetic alteration of individual cells to express high levels of either the VEGF receptor 2 or Notch showed that both stimuli caused endothelial cells to exit the cell cycle and increased their propensity to...
colonize arteries. To further investigate the downstream effects of Notch signalling on endothelial cell differentiation, an RNA-seq analysis was performed in mice with normal, diminished or increased Notch signalling. Notch-induced arterial endothelial cell differentiation was not dependent on upregulation of genes that determine arterial fate, but instead was dependent on the suppression of MYC-induced transcription of cell cycle and/or metabolism genes.

**Arterial expansion via cell recruitment**

The findings discussed above support a model in which arterial differentiation is coupled to cell cycle arrest (Fig. 1), which presents an intriguing question: how do arteries expand in size during development if their endothelial cell population does not proliferate? Numerous lineage-tracing studies from the past 5 years have shown that arteries grow by recruiting cells from veins and capillaries, rather than by expansion via endothelial cell division. In the mouse heart, timed labelling of Apj-expressing capillary plexus cells resulted in progressive pre-arterial specification and accumulation of endothelial cells in arteries during permissive developmental and postnatal time windows.

Three studies have assessed lineage-labelled arteries (BMXCreER), capillaries (Mfsd2aCreER) and veins (Gm5127(BAC)CreER) during postnatal retinal vasculature development in mice and showed that endothelial cells migrate in a vein-to-capillary-to-artery direction as the retinal vascular network expands and matures, but not in the opposite direction. Interestingly, these migration patterns are restricted to development and not seen in the mature eye. Furthermore, another study combined scRNA-seq analysis with endothelial cell lineage tracing and showed that endothelial cell migration from the vein to the artery is a common theme during development. The trajectory analysis performed on scRNA-seq data from mice at embryonic days 8–11 and matched human data predicted differentiation from vein to artery at multiple sites in the embryo in this early period. Computational predictions were validated by overlaying scRNA-seq data from lineage-labelled mouse vein endothelial cells (Nr2f2CreER). This propensity of endothelial cells to migrate into arteries might also explain why endocardium-derived vessels preferentially populate arterial segments in the outer wall of postnatal hearts.

**Endothelial cells migrate against blood flow**

The physical forces of blood flow drive endothelial cell migration towards arteries because endothelial cells are programmed to move against the direction of flow. Many questions remain regarding how endothelial cells are guided to perform this task, but several mechanisms have been proposed, including involvement of the bone morphogenetic protein (BMP) signalling pathway. Inhibiting the BMP receptor signalling components activin receptor type 1 (ALK1), endoglin or SMAD4 blocks blood flow-guided migration of endothelial cells in many settings. Loss of these components and subsequent loss of proper endothelial cell migration leads to arteriovenous malformations that form direct shunts between arteries and veins, bypassing capillaries. Zebrafish intersomitic vessel development is balanced by vein endothelial cell proliferation and polarization or migration against flow into arterial segments, a process that requires the actin cytoskeletal regulator Wasb (the homologue of human Wiskott–Aldrich syndrome protein) and flow-activated Notch.

Overactivation of endothelial cell migration against blood flow and into arteries has been achieved in mice by overexpressing Eng (encoding endoglin) or Dach1 (encoding dachshund homologue 1), the latter of which results in increased numbers of artery branches. Interestingly, loss of endothelial cell-derived non-canonical WNT5A and WNT11 caused endothelial cells to become over-responsive to flow, increasing their polarization and migration against flow. In the plexus, this augmented polarization and migration against flow increased pruning (the regression of a subset of microvessels within a growing vasculature), although the effect on arterial branching was not investigated. Further investigation into the mechanisms that drive
endothelial cell migration towards arteries for arterial expansion might provide the foundation for therapeutic applications in ischaemic diseases.

Compared with arterial endothelial cells, less is known about the development and maintenance of endothelial cells in veins and capillaries. Veins have transcriptional profiles that are distinct from those of arteries and the signals involved in inducing the venous cell state have mostly been studied in the early embryo. The establishment of veins requires cell autonomous endothelial cell phosphoinositide 3-kinase signalling and expression of COUP-TFI. Venous gene expression has been shown to be induced directly by BMP–BMP receptor type 1A (BMPRIA)–SMAD signalling in both mice and fish, ultimately leading to the formation of the receptor type 1A (BMPR1A)–SMAD signalling in both humans.

In capillaries in the brain and heart of both mice and humans, there is a unique barrier for the penetration of blood to Bowman’s space. The anatomical barrier for the penetration of blood to Bowman’s space is known as the blood–brain barrier (BBB) with tight intercellular junctions to restrict infiltration of blood into the brain. The endothelial cells of the brain form the blood–brain barrier unique to each organ.

Global insights from multi-organ studies

Bulk RNA-seq provided the first transcriptomic insights into the molecular heterogeneity of endothelial cells among different tissues, facilitating the identification of several tissue-specific endothelial markers for the heart, muscle, brain, liver and lung. ScRNA-seq corroborates the finding that endothelial cells from different tissues have specialized transcription programmes, which can be observed when endothelial cells in large multi-tissue datasets typically computationally separate (or cluster) into organ-specific subtypes. A detailed summary of scRNA-seq techniques that are commonly used in cardiovascular research is available elsewhere. Interestingly, organotypic endothelial cell specialization in mammals might be more exaggerated in blood endothelial cells because lymphatic endothelial cells cluster together regardless of the tissue of origin.

Single-cell analyses also facilitate the characterization of heterogeneity according to endothelial cell subtype (that is, artery, arteriole, capillary, venule and vein) (TABLES 2, 3). Endothelial cell heterogeneity across tissues is largely attributable to differential transcription in capillary endothelial cells; the endothelial cells in arteries and veins from different organs are more similar at the transcriptional level. Accordingly, ageing in the brain also affects capillary endothelial cells to a greater extent than arterial or venous endothelial cells.

The level of resolution afforded by single-cell analyses also establishes that organs with highly specialized vasculature, such as the brain, liver, lungs and kidneys, have more distinct transcriptomic profiles, whereas the gene expression of heart, skeletal muscle, diaphragm, mammary gland and adipose tissue tends to overlap. In addition, whereas endothelial cells from different organs share several major developmental pathways, including those relating to WNT signalling, cytokine regulation and metabolism, the expression patterns of the majority of genes associated with these pathways are unique to each organ.

Numerous studies have incorporated chromatin architecture information to gain further insights into the tissue-specific transcriptional regulation in endothelial cells, first by identifying DNA binding motifs in accessible chromatin regions and then comparing these motifs with the expression of genes encoding transcription factors to computationally predict endothelial transcription codes. Some consistent observations have emerged from these studies. Transcription factor motifs from the ETS transcription factor family, including transcriptional regulator ERG and Friend leukaemia integration 1 transcription factor, are universally enriched in endothelial cells. This finding is consistent with data showing that heterogeneous endothelial cell gene expression requires the combined function of both the endothelial cell subtype-specific transcription factors and ETS factors at enhancers. The source of heterogeneity in gene expression between tissues might stem from differences in distal, rather than proximal, cis-regulatory elements. In a study comparing assay for transposase-accessible chromatin using sequencing (ATAC-seq) peaks in endothelial cells from the liver, lung, brain and kidney, the majority of ATAC-seq
peaks at <2 kb from transcription start sites (proximal peaks) were shared between tissues, whereas the majority of ATAC-seq peaks at >2 kb from transcription start sites (distal peaks) were unique to individual tissues. Predicted tissue-specific endothelial cell transcriptional networks include lymphoid enhancer-binding factor or T cell factor family members in the brain, which are well-known to be downstream effectors of canonical WNT signalling. GATA transcription factors seem to regulate heart-specific and liver-specific endothelial

| Organ system | Species | Developmental stage | Dataset contents | Single-cell capture and scRNA-seq method | Refs. |
|--------------|---------|---------------------|------------------|------------------------------------------|-------|
| Multi-organ  | Mouse   | Adult               | scRNA-seq        | Smart-seq2 and 10x Genomics GemCode       | 33    |
|              |         |                     | scRNA-seq        | Microwell-seq                            | 73    |
|              |         |                     | scRNA-seq, RNA-seq, ATAC-seq, MethylC-seq | 10x Genomics GemCode                      | 33    |
|              | Human   | Embryonic (E72–E129) | sci-ATAC-seq     | sci-ATAC-seq                              | 75    |
|              |         | Adult (22–74 years)  | scRNA-seq        | Smart-seq2 and 10x Genomics GemCode       | 33    |
|              | Heart   | Embryonic (E12.5–E14.5) | scRNA-seq        | 10x Genomics GemCode                      | 33    |
|              | Mouse, human | Embryonic mouse (E12, E17.5), embryonic human (11 weeks, 14 weeks, 22 weeks), adult mouse | scRNA-seq | 10x Genomics GemCode                      | 33    |
|              | Human   | Embryonic (5–25 weeks) | scRNA-seq        | STRT-seq                                  | 77    |
|              |         | Adult (40–75 years)  | scRNA-seq        | 10x Genomics GemCode                      | 33    |
|              | Lung    | Embryonic (E12.5–E17.5), postnatal (P3–P42) | scRNA-seq, scATAC-seq | 10x Genomics GemCode                      | 33    |
|              | Postnatal (P1) | scRNA-seq                              | Drop-seq                  | 80    |
|              | Postnatal (P3, P7, P14) | scRNA-seq                              | MULTI-seq                 | 81    |
|              | Human   | Adult               | scRNA-seq        | Smart-seq2 and 10x Genomics GemCode       | 33    |
|              |         | Adult               | scRNA-seq        | 10x Genomics GemCode                      | 33    |
|              | Brain   | Embryonic (E14.5)   | RNA-seq (TRAP-seq), scRNA-seq | TruSeq v2                                | 84    |
|              |         | Adult               | scRNA-seq        | Smart-seq2                                | 67    |
|              | Mouse, human | Adult mouse, paediatric and adult human | snRNA-seq      | 10x Genomics GemCode                      | 67    |
|              | Human   | Fetal, adult        | scRNA-seq        | 10x Genomics GemCode                      | 67    |
|              |         | Adult               | scRNA-seq, RNA-seq | 10x Genomics GemCode                      | 67    |
|              | Kidney  | Embryonic           | scRNA-seq        | 10x Genomics GemCode                      | 67    |
|              | Mouse   | Embryonic E14.5     | scRNA-seq        | 10x Genomics GemCode                      | 67    |
|              | Embryonic E14.5 | scRNA-seq              | Drop-seq, Chromium 10x Genomics and Fluidigm C1 | 67    |
|              | Embryonic, adult | snRNA-seq, sn-ATAC-seq | Smart-seq2 and 10x Genomics GemCode       | 67    |
|              | Human   | Embryonic, paediatric, adult | scRNA-seq   | 10x Genomics GemCode                      | 67    |
|              | Embryonic (12–18 weeks) | scRNA-seq                  | Drop-seq                  | 67    |
|              | Embryonic (9–19 weeks) | scRNA-seq                  | 10x Genomics GemCode       | 67    |
|              | Adult   | snRNA-seq, sn-ATAC-seq | 10x Genomics GemCode       | 67    |
|              | Liver   | Embryonic (E12–E18), postnatal (P2–P30) | scRNA-seq, bulk RNA-seq | 10x Genomics GemCode                      | 67    |
|              |         | Adult               | scRNA-seq        | MARS-seq                                  | 67    |
|              |         | scRNA-seq           | 10x Genomics GemCode       | 67    |
|              |         | scRNA-seq           | 10x Genomics GemCode       | 67    |
|              | Human   | Adult               | scRNA-seq         | CEL-seq2                                  | 67    |
|              |         | scRNA-seq           | 10x Genomics GemCode       | 67    |

ATAC-seq, assay for transposase-accessible chromatin using sequencing; CEL-seq2, cell expression by linear amplification and sequencing; Drop-seq, droplet sequencing; E, embryonic day; MARS-seq, massively parallel RNA single-cell sequencing; MULTI-seq, multiplexing using lipid-tagged indices for single-cell and single-nucleus RNA sequencing; P, postnatal day; scRNA-seq, single-cell RNA sequencing; snRNA-seq, single-nucleus RNA sequencing; sci-ATAC-seq, single-cell combinatorial indexing assay for transposase-accessible chromatin using sequencing; sci-RNA-seq, single-cell combinatorial indexing RNA sequencing; Smart-seq2, switching mechanism at the 5′ end of the RNA template sequencing; sn-ATAC-seq, single-nucleus combinatorial indexing assay for transposase-accessible chromatin using sequencing; STRT-seq, single-cell tagged reverse transcription sequencing; TRAF-seq, targeted purification of polysomal mRNA.
cell transcription, whereas homeobox transcription factor family members are enriched in the lungs and kidneys33,109.

Multi-organ datasets have provided support for earlier observations that surrounding cells induce tissue-specific behaviour in endothelial cells, whereas endothelial cells in turn affect the transcriptomic profiles of their neighbours. For example, subpopulations of cardiac endothelial cells express cardiac muscle-specific genes and sinusoidal endothelial cells have similar gene expression profiles to those of adjacent hepatocytes29,98. Numerous studies have begun to investigate the functional implications of these interactions across a number of organ systems. A study combining scRNA-seq analysis with bone marrow transplantation experiments characterized the crucial role of arterial endothelial cells in establishing haematopoietic stem cell colonization of the bone marrow12,13. Another study similarly used scRNA-seq to probe the complex response of endothelial cells to pericyte loss in the brain, a phenomenon seen in a wide range of neurological diseases14.

Finally, sexual dimorphisms are known to contribute to differences in organ function, susceptibility to disease and ageing, processes that involve blood vessels that permeate almost every tissue115. Transcriptomic and proteomic analyses have begun to establish that these differences, at least in the heart, are attributable to both hormonal control and sex chromosome composition115. Many organism-wide scRNA-seq analyses control for the possibility of sex-dependent differences in gene expression by utilizing tissues from a single sex, but only a few studies have assessed the effects of sex on organ-specific

Table 2 | Representative selection of marker genes for mouse organ-specific vascular subtypes

| Organ system       | Developmental stage | Vascular EC subtype | Marker genes | Refs. |
|--------------------|---------------------|---------------------|--------------|-------|
| Multi-organ        | Embryonic or adult  | Pan-endothelial      | Pecam1, Kdr, Cldn5, Emsn, Cdhn5, Tie1, Egf17 | 2,80,85,113 |
|                    |                     | Vein                | Apj, Nr2f2, Vwf | 2005,05 |
|                    |                     | Artery              | Gja4, Mecom  | 2005 |
| Heart              | Embryonic           | Artery              | Gja4, Gja5  | 65 |
|                    | Pre-artery or arterio|                    | Gja4, Gja5, Unc5b, Hey1, Slc45a4 | 2005 |
|                    | Vein                |                     | Nr2f2, Apj, Nr2f2 | 2005 |
|                    | Capillary           | Tissue-specific EC  | Wt1, Slc28a2, Eepd1, Kcnq5, Carp8, Fbn1, Meox2, Rnf1, Lamb1, Mmyd16 | 2 |
|                    |                     | Artery              | Apnl, Dil4, Notch1, Mecom, Igfbp3, Cxcr4, Cx40 | 29 |
| Lung               | Adult               | Artery              | Map, Cdh13, Htral1, Bmx, Fbn2, Sulf1 | 131 |
|                    |                     | Vein                | Slc6a2, Bst1, Car8, Amig2, Mutn1, Csrp2 | 131 |
|                    | Tissue-specific EC  | Grtp1, Adrb1, Scl7a, Tmem100, Hpgd, Foxf1a, Ncknp5, Rasgef1a, Fenadr, Fnx | 2 |
|                    | Capillary (aCap)    | Car4, Ednrb, Fbn1, Tbx2, Rprml, Chat1, Apnl | 82,131 |
|                    | Capillary (gCap)    | Apnl, Gpihp1, Pvlap, Cdh93, Ptpbr, Cemip2, Tek, Cxcl12 | 82,131 |
| Brain              | Embryonic           | Tissue-specific EC  | Foxf2, Foxl2, Foxq1, Lef1, Ppard, Zfp551, Zic3 | 84 |
|                    | Adult               | Tissue-specific EC  | Slco1c1, Slco1a4, Slc22a8, Mfsd2a, Slc38a3, Spock2, Foxf2, Edn3, Sra8, Slc38ak | 2 |
|                    | Artery              | Gkn3, Sema3g, Efnb2 | 65 |
|                    | Venule              | Slc38a5             | 65 |
|                    | Capillary or vein   | Slc16a1             | 65 |
| Kidney             | Adult               | Tissue-specific EC  | Egf17, Drum1, Dkk2, Esm1, Igfbp5, Pbx1, Boc, Igfbp3, Irg3, Tfjnpl2, Ptptru | 87,88 |
|                    | Glomerular EC       | Pi16, Plat, Ehd3, Cyb4b1, Tspan7, Lpl | 143 |
|                    | Cortical EC         | Igfbp3, Pvlap, Npr3 | 143 |
|                    | Medullary EC        | Igf1, Cryab, Igfbp7, Cd36, Aap1, Ifij712a | 143 |
| Liver              | Embryonic or adult  | Portal vein         | Ly6a, Cd34, Cd9, Ephb2, Gja5, Sox17 | 97 |
|                    | LSEC                | Mrc1, Fcgr2b, Stab1, Stab2 | 97 |
|                    | Adult               | Tissue-specific EC  | c-Maf, Clec4g, Fcgr2b, Stab2, Oit3, Bmp2, Aass, Mrc1, Plxnc1, Wnt2 | 2,136 |
|                    | LSEC, zone 1        | Dil4, Efnb2, Ltbp4, Msr1, Ntrn4 | 98,99 |
|                    | LSEC, zone 2 or 3   | Rspo3, Wnt2, Wnt9b | 98,99 |

aCap, aerocyte capillary; EC, endothelial cell; gCap, general capillary; LSEC, liver sinusoidal endothelial cell.
transcriptomic profiles\(^{106}\). Using the Tabula Muris data-set, sex was found to be an important factor underlying endothelial cell transcriptome heterogeneity, but only in some tissues, including the aorta, brain and lungs\(^ {107}\). Future cell atlas studies should incorporate sex as a parameter to better understand its influence on vascular physiology.

**Organ-specific findings at high resolution**

**Cardiac vasculature.** The majority of coronary blood vessels develop via convergent differentiation of endothelial cells from two distinct progenitor sources: the inlet vein known as the sinus venosus and the endocardium that lines the heart chamber\(^ {42,116–120}\). Cells from these two populations ultimately localize to complementary yet overlapping regions of the heart. The sinus venosus mostly forms vessels in the outer ventricular walls and the endocardium mostly forms vessels in the septum and inner ventricular wall. Despite this spatial divergence, endothelial cells from both progenitor sources begin to sprout between embryonic days 11 and 12.5 in mice\(^ {116}\). The sinus venosus and endocardium initially sprout in response to distinct molecular signals, specifically epicardial VEGFC–apelin receptor early endogenous ligand signalling and SMAD signalling for sinus venosus–derived cells, and intramyocardial

### Table 3 | Representative selection of marker genes for human organ-specific vascular subtypes

| Organ system | Developmental stage | Vascular EC subtype | Marker genes | Refs. |
|--------------|---------------------|---------------------|--------------|-------|
| Multi-organ  | Embryonic or adult  | Pan-endothelial      | PECAM1, CDH5, VWF, KDR, FLT1, TEK, CLDN5 | 68,70,82,83,85 |
|              |                     | Artery              | GJA4, GJA5, HEY1, GATA2, CXCRR4, SOX17, MECOM | 96,85,87,90 |
|              |                     | Vein                | ACKR1, NR2F2, PLVAP | 68,80,75 |
| Heart        | Embryonic           | Artery              | GABBR2, GRIA2, SSUH2, JAG1 | 66 |
|              |                     | Vein                | ACKR1, LHX6, SELE | 66 |
|              |                     | Capillary           | PRDM1, INMT, APLNR, CA4 | 66 |
|              | Adult               | Tissue-specific EC  | SLC14A1 | 91 |
|              |                     | Artery              | SEMA3G, EFNB2, DLL4 | 78 |
|              |                     | Vein                | NR2F2, ACR1 | 78 |
|              |                     | Capillary           | RGCC, CA4 | 78 |
| Lung         | Adult               | Tissue-specific EC  | VIPR1 | 91 |
|              |                     | Artery              | DKK2, SERPINE2 | 131 |
|              |                     | Vein                | CPE, PTGDS, C7, PLA1A | 131 |
|              |                     | Bronchial EC        | VWA1, HSPG2, PLVAP, MYC, HBEGF | 82 |
|              |                     | Capillary (aCap)    | ADIRF, S100A4, EMD, EDNRB, SOSTDC1, ILIRL1, APLN | 131 |
|              |                     | Capillary (gCap)    | FCN3, EDN1, SLC6A4, GPHBP1, CD36, IL7R, VWF, PTPRB, PLVAP | 131 |
| Brain        | Adult               | Large artery        | LTBP4 | 69 |
|              |                     | Artery              | INTS6, HSPA1A, JUNB, MECON, ARL15, TXNIP, MGP, ADAMTS1 | 68,69 |
|              |                     | Arteriole           | VEGFC, ARL15, BMX, EFNB2, WISR, AIF1L, CD320 | 67 |
|              |                     | Venule              | TSHZ2, ADGRG6, SLC38A5, LRRC1, BNC2, ETF6, TEMEM132C, ATP10A, TAM2, PRCP, PRSS23, RAMP3 | 67–69 |
|              |                     | Vein                | ACKR1, ILIR1, TSHZ2, PTGDS, POSTN, DNASE1 | 65,69 |
|              |                     | Large vein          | CCL2 | 69 |
|              |                     | Capillary           | MFSD2A, SLC7A5, TFRC, SLC38A5, SRARP, RGCC, SLC3A2, BSG, SLC16A1, SLCO1A2 | 67–69 |
| Kidney       | Embryonic           | Tissue-specific EC  | NOTCH4 | 90 |
|              | Paediatric or adult | Glomerular EC       | SEMA3G, CLDN5 | 89 |
| Liver        | Adult               | Tissue-specific EC  | OIT3 | 91 |
|              |                     | Portal vein         | AQP1, CD9, IFITM1, RAMP3, INMT, DNASE1L3, LIFR | 97,101 |
|              |                     | LSEC                | CLEC4G, STAB1, STAB2, CD14 | 97 |
|              |                     | LSEC zone 1         | MGP, SPARCL1, TM4SF1, CLEC14A, BTNL9, ANPEP | 100,101 |
|              |                     | LSEC zone 2 or 3    | CCL14, CLEC1B, FCN2, S100A13, LYVE1, FCN3 | 100,101 |
|              |                     | Central vein        | SELP, SELE, VWF | 97 |

\(a\)Cap, aeryocyte capillary; EC, endothelial cell; gCap, general capillary; LSEC, liver sinusoidal endothelial cell.
VEGFA for endocardium-derived cells. After sprouting onto the developing heart, previously fully differentiated sinus venosus vein endothelial cells or endocardium will undergo extensive cell state changes within the coronary plexus and remodel into coronary arteries, capillaries and veins.

After birth, neonatal cardiac ventricles undergo rapid growth and remodelling. The remaining trabecular myocardium, which lacks intramyocardial vessels during embryogenesis, undergoes compaction and acquires rich coronary vascularization. Endocardium-derived vessels have been shown to expand into this region in the weeks after birth, and new vessels have been suggested to arise from postnatal lineage conversion of endocardial cells to coronary vessel endothelial cells. Stage-specific lineage-tracing experiments have further refined this model by showing that late fetal and neonatal endocardial cells contribute minimally to the formation of postnatal coronary vessels, even with injury. Differentiation of endocardial cells into coronary endothelial cells occurs mostly during early heart development (before embryonic day 13.5 in mice) and endocardium-derived vascular tunnels do not adopt a mature artery fate until postnatal day 7 in mice. Instead of endocardial conversion, postnatal endothelial cells from coronary vessels in the inner myocardium express high levels of VEGFR3 and undergo angiogenic expansion in a delta-like protein 4–NOTCH 1-dependent manner to vascularize the newly compacted myocardium. This process is mediated by capillary endothelial cells from other organs, which adopt a ‘convergent differentiation’ has also been seen in olfactory projection neurons, but not in certain populations of haematopoietic cells or tissue-resident macrophages, which retain markers of primed fate potential. Furthermore, no functional differences have been observed in endothelial cells from either lineage, such that they have largely identical proliferative capacity in response to injury.

Techniques such as reference mapping, as well as comparisons of mouse and human scRNA-seq datasets through robust integration, provide information about the transcriptional similarities and differences between species. These analyses of developing mouse and human hearts revealed that the developmental environment and endothelial cell subtype transcriptional states are similar in both species, validating the use of mice as a model for many aspects of human coronary development. Follow-up functional experiments used these data in an induced pluripotent stem cell culture model to determine that the histone–lysine N-methyltransferase MECOM is crucial for human artery endothelial cell differentiation. Furthermore, trajectory analyses of human data predicted a transitory cell population from the endocardium, which had a gene expression pattern validated using lineage tracing that overlapped with that of the transitory cell population in mice. Therefore, despite the dearth of lineage-tracing techniques for human studies, overlapping human and mouse datasets provide some evidence that coronary vessels from humans also arise from the endocardium.

Lung vasculature. Alveoli, the terminal airspaces in the mammalian lung and a major site of gas exchange, are surrounded by a dense network of capillaries. Single-cell studies have identified heterogeneities in alveolar vasculature that are crucial for structural integrity. Alveolar development during late embryogenesis is accompanied by the emergence of a specialized capillary subtype called CAR4+ aerocyte capillaries (aCaps), which intermingle with the ‘general capillaries’ (gCaps).

CAR4 encodes carbonic anhydrase 4, which catalyses the reversible hydration of carbon dioxide into bicarbonate and protons. Although gCaps have similar gene markers to capillary endothelial cells from other organs, aCaps express markers such as ICAM1 (encoding intercellular adhesion molecule 1) and EDNRB (encoding endothelin receptor type B), and are unique to the lung. aCaps are also atypically large with a ‘Swiss cheese’-like morphology, spreading over the thin alveolar type 1 epithelium where gas exchange occurs. This location and their gene expression patterns indicate that aCaps have specialized roles in gas exchange and leukocyte trafficking.

Trajectory analysis of scRNA-seq and lineage-tracing data revealed that gCaps are progenitors for aCaps during alveolar development and after injury in adult[13]. Development and maintenance of aCaps specifically require VEGF from airway epithelium, given that VEGF deletion results in depleted aCap numbers but does not affect gCap numbers. aCaps also seem to be required for lung maturation because lack of aCaps in mice results in enlarged alveoli despite the presence of myofibroblasts.

Together, this deeper understanding of capillary heterogeneity has informed human pathology. As in the heart, interspecies comparisons have revealed that all endothelial cell subtypes found in mice are present in humans, and that CAR4+ aCaps also emerge during late embryogenesis. Analysis of a human lung adenocarcinoma sample revealed a loss of aCaps and gain of a transcriptionally intermediate population that co-expressed markers of aCaps and gCaps. This same phenomenon was observed in single-cell atlases of lungs from patients who died of coronavirus disease 2019 (COVID-19) (Fig. 2b). Interestingly, endothelial cells were the second most abundant site of viral RNA load in these patients, despite low-to-absent expression of the entry receptor for SARS-CoV-2 [REF.], consistent with the
knowledge that endothelial cells are a major site of pathology in patients with COVID-19 (Ref. 136). A comparison of hyperoxia-induced bronchopulmonary dysplasia in mice with matched human samples showed that aCaps mediate the increased expression of INHBA, which encodes inhibin-β A chain 81, a member of the TGFβ superfamily that has been suggested to contribute to bronchopulmonary dysplasia. Furthermore, the brain also shows conserved endothelial cell populations and zonation between mice and humans and a greater diversity of human perivascular cells. Two transcriptionally distinct populations of capillary endothelial cells exist in the lung: general capillaries (gCap) and aerocyte capillaries (aCap). These two capillary populations have different roles in lung maturation and mature lung function. In disease states such as adenocarcinoma, a transcriptional intermediate between gCap and aCap arises in both humans and mice. This transcriptional intermediate has also been seen in the lungs of patients with coronavirus disease 2019 (COVID-19). Overlapping features of inflammation and cellular stress can also be seen in both capillary subtypes in advanced chronic obstructive pulmonary disease. AV1, alveolar type 1 cell; RBC, red blood cell; t-SNE, t-distributed stochastic neighbour embedding.

Brain vasculature. Given the prominent role of the central nervous system blood vessels in human disease and ageing 30, these vessels have been the subject of many single-cell transcriptomic studies in the past 5 years. Indeed, single-cell analyses of brain-specific capillaries led to the identification of a previously unappreciated feature of many capillary beds. Capillaries comprise the vast majority of blood vessels and are the main site of oxygen exchange, yet multi-organ studies have thus far failed to identify a unifying capillary molecular marker 3. This lack of a unifying capillary marker might be attributable to the observation that instead of being a homogeneous entity, cells within a single capillary bed exist along an arterial–venous transcriptional continuum, referred to as zonation, which depicts gradual cellular phenotypic changes along an anatomical axis (Fig. 1). This feature of capillaries was first confirmed in the brain (via an approach that assessed and visualized data using distance matrices) and later in the heart (using artery or vein gene scores) 29,65. Interestingly, zonation is not a phenotype shared by endothelial cell-adjacent mural cells in the central nervous system vasculature. Arteriole smooth muscle cells abruptly transition to become pericytes at the arteriole–capillary boundary in both mice and humans 29,65.

Fig. 2 | scRNA-seq reveals novel endothelial cell types. a | Comparison of human and mouse single-cell RNA sequencing (scRNA-seq) datasets for heart and brain vasculature reveals similar endothelial cell states between species. Although expression of unique genes exists for endothelial cell subtypes within each species, these analyses support the use of mouse models for studying human development and pathology. Key findings from these datasets reveal that the heart has conserved endothelial cell populations and similar capillary endothelial cell states in mice and humans and unique human artery specification genes. Furthermore, the brain also shows conserved endothelial cell populations and zonation between mice and humans and a greater diversity of human perivascular cells. b | Two transcriptionally distinct populations of capillary endothelial cells exist in the lung: general capillaries (gCap) and aerocyte capillaries (aCap). These two capillary populations have different roles in lung maturation and mature lung function. In disease states such as adenocarcinoma, a transcriptional intermediate between gCap and aCap arises in both humans and mice. This transcriptional intermediate has also been seen in the lungs of patients with coronavirus disease 2019 (COVID-19). Overlapping features of inflammation and cellular stress can also be seen in both capillary subtypes in advanced chronic obstructive pulmonary disease. AV1, alveolar type 1 cell; RBC, red blood cell; t-SNE, t-distributed stochastic neighbour embedding.
Other studies have also identified capillary zonation in human brains, early mouse and human embryos and developing human hearts\(^\text{[26,69-72]}\) (Fig. 2a). Loss of communication between pericytes — a cell type reported to regulate the formation of the BBB\(^\text{[72,74]}\) — and endothelial cells was also found to skew capillary zonation towards a venous identity rather than causing widespread loss of BBB function and ectopic tips\(^\text{[13]}\).

Finally, single-cell analyses can also provide insights into the source of disease that results from genetic variants. Findings from the cross-referencing of cell subtype gene expression in mice with gene variants associated with Alzheimer disease or genes with altered expression in brain tissue from patients with Alzheimer disease suggest that capillary endothelial cells are a potential source of pathology, possibly through changes in BBB-related genes\(^\text{[10]}\). The same analysis using human data pinpointed endothelial cells and mural cells as potential sites of disease\(^\text{[9]}\).

**Kidney vasculature.** Given that the kidney has diverse and distinctive endothelial cell populations, the renal vasculature is a good candidate for single-cell analysis\(^\text{[14,15]}\). The renal microvascularity regulates blood flow, facilitates filtration, modulates inflammation and maintains physiological blood pressure. These functions are compartmentalized to specific regions: glomerular endothelial cells make up the filtration barrier, whereas tubular and peritubular capillary endothelial cells absorb and secrete fluids and other substances\(^\text{[6,14]}\). Single-cell analyses of mouse models of kidney disease have provided insights into the functional divisions between cortical, medullary and glomerular endothelial cells and identified cell type-specific responses to pathology. Cortical endothelial cells express high levels of *Igfbp3* (encoding insulin-like growth factor binding protein 3) and *Npr3* (encoding natriuretic peptide receptor 3) among other genes, whereas medullary endothelial cells express *Igf1* (encoding insulin-like growth factor 1) and *Cd36* (encoding platelet glycoprotein 4). Glomerular endothelial cell-enriched genes include *Ehd3* (encoding EH domain-containing protein 3), *Cyp4b1* (encoding cytochrome P450 4B1) and *Tspan7* (encoding tetraspanin 7)\(^\text{[145-147]}\) (Fig. 5a). In response to acute water deprivation, medullary endothelial cells show the greatest transcriptional changes, upregulating the expression of genes related to oxidative phosphorylation in an effort to promote survival and concentrate urine\(^\text{[145]}\). A mouse model of nephritis (via nephrotoxic antibody—endothelial cell pairs back to their expression-inferred tissue coordinates\(^\text{[96]}\). Molecular signatures for subpopulations of pericentral liver endothelial cells were identified and the zonation of endothelial cells across lobules was characterized\(^\text{[98]}\). An analysis of the spatially resolved proteome revealed a high level of concordance with the transcriptomic data. Interestingly, although zonation in the brain corresponds to a continuum of arterial–venous transcriptional states, several studies have found that zonation in the liver relates to functional differences across the lobule axis\(^\text{[98,101]}\).

Endothelial cell zonation was found to drive much of this functional zonation in hepatocytes via a TIE–WNT signalling axis\(^\text{[150]}\). The initial establishment of zonation in embryonic endothelial cell progenitors depends on MAF, a transcription factor crucial for sinusoid (or liver capillary) identity\(^\text{[97]}\).

As with capillaries in the kidney, liver sinusoidal endothelial cells have zone-specific susceptibility to injury inflicted during disease, such as sinusoidal endothelial cells in lobule zone 3 that are particularly affected in cirrhotic liver of mice\(^\text{[151]}\). Just as aCaps and gCaps in the lung lose their specialized transcriptional states in disease states, cirrhosis causes zone 3 sinusoidal endothelial cells to lose their specialized fenestrae through a process known as capillarization\(^\text{[152]}\).

**Advances in organoid vasculature**

Recent developments in single-cell genomics have revolutionized our understanding of the heterogeneity of vascular cells and their exquisitely regulated development within tissue microenvironments. As well as facilitating the identification of rare and novel endothelial cell types, such as specialized capillary subtypes in the lungs, liver, kidneys and retina, scRNA-seq has offered a higher resolution window into the transcriptional changes that occur during disease\(^\text{[92,93,148]}\). Furthermore, scRNA-seq data have provided new insights into the interactions between endothelial cells and adjacent cell types, such as smooth muscle cells, pericytes and immune cells, and the roles that these interactions have in both the development and the maintenance of organ-specific vasculature. These findings, combined with results from sophisticated lineage-tracing studies, raise important considerations for tissue engineering. A lack of physiologically accurate perfusable vasculature is one of the most important challenges limiting the expansion of multicellular ‘mini-organs’ known as organoids\(^\text{[144,146]}\). Advances in differentiation protocols and culture techniques for human pluripotent stem cells (hPSCs)\(^\text{[153]}\) have resulted in the development of 3D self-organizing human blood vessel organoids, which are capable of forming complex vascular networks that recapitulate many of the mural cell–endothelial cell interactions known to be crucial for supporting structures such as the BBB\(^\text{[154]}\). These organoids, as well as...
several hPSC-derived liver\textsuperscript{154,155}, kidney\textsuperscript{156} and brain\textsuperscript{157} organoids, have shown promising, though often partial, levels of host vascularization in vivo after transplantation into immunodeficient mice.

Although in vivo transplants are an exciting demonstration of the therapeutic potential of organoids, the generation of endogenous organ-specific vasculature remains crucial. By using scRNA-seq data, one group demonstrated that hPSC-derived intestinal organoids are capable of inducing the differentiation of endothelial cells with an intestinal-specific transcriptional signature\textsuperscript{158}. Other efforts have focused on increasing...
the complexity and functionality of organ-specific vasculature by modulating key signalling pathways and exposing in vitro systems to physiologically relevant physical forces. Through a combination of these methods, several groups have developed increasingly sophisticated cardiac organoids complete with internal lumens and vascular networks. Other groups have shown that modulating morphogen gradients and fluid shear stress conditions using microfluidic systems improves the expansion of endothelial progenitors in the kidney, although further analysis is required to determine whether faithful endothelial cell subtype differentiation has been achieved with these methods.

Finally, 3D bioprinting offers an alternative approach to embedding perfusable vascular channels for larger regions of tissue. In a newly developed biomaterializing method, multiple hPSC-derived organoids, embryoid bodies or spheroids can be combined into a granular tissue matrix. Sacrificial ink is injected and then removed to create channels within the tissue that can be seeded with endothelial cells. This approach has resulted in the successful vascularization of a contractile cardiac organoid matrix and might be capable of producing more structurally accurate organ-specific vasculature than other methods.

Conclusions
In this Review, we discuss the current molecular understanding of the differentiation of endothelial progenitors into arteries and veins. New insights are used to form a model in which artery differentiation is coupled to cell cycle arrest, whereas vein differentiation is associated with cell proliferation and direct induction of the BMP-BMPRIA-SMAD pathway. Arteries then expand by recruiting cells from capillaries and veins, which migrate against the direction of blood flow to expand the arterial network. Numerous multi-organ scRNA-seq and single-cell multi-omic atlases developed in the past 5 years have led to the discovery of novel endothelial cell-specific findings for several critical cardiovascular organs. Capillary endothelial cells are the primary source of vascular heterogeneity across organs, and findings from the past 5 years have shown that they are also the primary source of transcriptional changes in multiple disease states. A general trend observed in the liver, lungs and brain is that disease promotes a loss of specialized capillary function and a reversal to a more general transcriptional state. These findings have implications for the development of targeted therapies and for the differentiation of improved organ-specific vasculature for tissue-engineering applications.

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