Endothelial struts enable the generation of large lumenized blood vessels de novo

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De novo blood vessel formation occurs through coalescence of endothelial cells (ECs) into a cord-like structure, followed by lumenization either through cell–1–3 or cord-hollowing4–7. Vessels generated in this manner are restricted in diameter to one or two ECs, and these models fail to explain how vasculogenesis can form large-diameter vessels. Here, we describe a model for large vessel formation that does not require a cord-like structure or a hollowing step. In this model, ECs coalesce into a network of struts in the future lumen of the vessel, a process dependent upon bone morphogenetic protein signalling. The vessel wall forms around this network and consists initially of only a few patches of ECs. To withstand external forces and to maintain the shape of the vessel, strut formation traps erythrocytes into compartments to form a rigid structure. Struts gradually prune and ECs from struts migrate into and become part of the vessel wall. Experimental severing of struts resulted in vessel collapse, disturbed blood flow and remodelling defects, demonstrating that struts enable the patency of large vessels during their formation.

Cord-hollowing during vasculogenesis can either occur through fusion of intracellular lumens (formed through pinocytosis) of endothelial cells (ECs) connected in a head-to-tail orientation1–3 or through the formation of an extracellular lumen between two ECs. This latter model depends on apical–basal polarization of ECs, which leads to the rearrangement of cell–cell junctions to the periphery of two adjacent ECs. Delivery of negatively charged glycoproteins to the apical side of the cells subsequently leads to the formation of a lumen between two ECs through electrostatic repulsion2–4. Alternatively, extracellular lumen could also result from apical exocytosis of vacuoles4. As a result, the diameter of these vessels is restricted to either one (intracellular lumen) or two ECs (extracellular lumen). However, because vasculogenesis can result in the formation of large vessels, it raises the question of how these vessels acquire their large diameters. A good example of such a large-diameter vessel is the posterior cardinal vein (PCV), which forms along with the dorsal aorta (DA) the first blood vessels in the developing vertebrate embryo. In zebrafish, the caudal portion of the PCV, posterior to the yolk sac extension, is referred to as the caudal vein (CV) and is approximately five times larger than the DA and thus the largest blood vessel observed during development. Despite this large difference in diameter, the DA and PCV are formed and become functional over a nearly equivalent time-frame, suggesting that the lumen of the CV is probably not formed by one of the currently known mechanisms. To investigate how the CV is formed, we visualized its formation from 18 hours post fertilization (h.p.f.) onwards, the time point when ECs start to form the DA and CV at the midline of the embryo5. For visualization purposes, we focused either on the area of lumen formation (Fig. 1a, orange box) or vessel wall formation (Fig. 1a, green box). Around 18 h.p.f., the most medially positioned ECs started to coalesce into a network spanning the future lumen of the CV (Fig. 1b, orange boxes and Supplementary Video 1, left panel). Most branches of this network are composed of multiple ECs (Extended Data Fig. 1a). By time-lapse imaging, we found that this network gradually prunes and disappears from the lumen by 26–28 h.p.f. (Fig. 1b, orange boxes, 1c (blue area), Extended Data Fig. 1b and Supplementary Video 1, left panel). The wall of the CV is initially composed of only a few patches of ECs (18 h.p.f.), in stark contrast with the current models of vasculogenesis, in which the vascular cord already forms a ‘leak-proof’ vessel before lumenization (Fig. 1b, green boxes and Supplementary Video 1, right panel). Given that closure of the vessel wall coincided with pruning of the endothelial network, we hypothesized that ECs from the network are reused to form the CV wall. To test this hypothesis, we traced ECs within individual branches of the network by ultraviolet photoconversion of the DENDRA2 fluorescent protein from green to red. We found that the majority of the converted ECs (in 17 of 25 embryos) had integrated into the wall of the CV 20 h after conversion (Fig. 1d). In roughly 30% of the cases, we found that converted cells integrate into the arterial system, with ECs integrated into the floor of the DA or as single cells in the sub-aortic space between the DA and PCV, a typical appearance of haematopoietic stem and progenitor cells (Fig. 1c)5,9–10. Thus, the CV is formed around a temporary network of both arterial and venous ECs, which seems to maintain the patency of the CV. Hence, we termed these structures endothelial struts. Of note, we also found struts in the anterior part of the PCV, consisting only of 1–2 ECs, consistent with the smaller diameter relative to the posterior CV (Extended Data Fig. 1c). Furthermore, we did not find any evidence of EC proliferation or apoptosis during the formation or pruning of struts, suggesting that, through an intricate mechanism of ECs coalescing into struts, pruning and EC migration, large-diameter blood vessels can be formed without the need to expand the initial pool of ECs (Extended Data Fig. 1d).

Our photoconversion experiments suggested that struts in the CV contained arterial ECs (Fig. 1c). This finding is intriguing, because arterial and venous precursors arise from two distinct locations

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in the lateral plate mesoderm and were hypothesized to form the DA and PCV as independent entities\(^1\). Our images from the trunk at 20 h.p.f. showed two separate populations in the anterior part of the trunk that become increasingly intermingled, without a clear separation in the posterior part of the trunk (CV region; Fig. 2a). The arterial and venous ECs in the CV region formed a common precursor vessel that, over time, unmixed into the DA and CV in an anteroposterior direction (Fig. 2b and Supplementary Video 2). We visualized arterial and venous ECs by imaging embryos in which arterial ECs were fluorescently labelled by the arterial-restricted Notch ligand \(d\ell4\) or by fluorescent reporting of Notch activity (\(\text{Tp}1\)) (Fig. 2c,d). Consistent with results from our photoconversion experiments, we found that some ECs in struts located within the CV expressed \(d\ell4\) or were positive for Notch activity (\(\text{Tp}1\)) (Fig. 2c,d). Of note, we observed that the majority of arterial ECs that were juxtaposed to the hypochord, forming the roof of the DA, had a higher GFP signal (\(d\ell4\)) and showed increased Notch activity (\(\text{Tp}1\)) compared to arterial ECs positioned more ventrally (Fig. 2c,d). These findings suggest that arterial ECs positioned against the hypochord have a stronger arterial signature than the more ventrally positioned arterial ECs, which include those forming endothelial struts. From this, we reasoned that unmixing of arterial and venous ECs in the CV occurs through the gradual increase of arterial genes in the more ventrally positioned arterial ECs, including struts. To test this, we inhibited arterial differentiation by abrogating Notch signalling through administration of
Fig. 2 | The caudal vein and dorsal aorta arise from a common precursor vessel through unmixing of arterial and venous ECs. a, Arterial and venous EC populations are indicated by anatomical position. The right panel shows pseudo-coloured populations. The images are representative of 20 analysed embryos from two independent experiments. b, Stills from Supplementary Video 2. The CV and DA are formed through segregation of arterial and venous ECs from a common precursor vessel. Red colour and arrows depict the formation of the DA lumen. The stills are representative of five independent time-lapse experiments in which one embryo is imaged each. c, Arterial ECs are marked by GFP (dll4; red arrowheads), all ECs are marked by laCherry (magenta) and the white arrowhead depicts the strut. The images are representative of 20 analysed embryos from two independent experiments. d, Notch signalling is reported by the expression of eGFP under the control of 12xCSL Notch-responsive elements (Tp1, red arrowheads), all ECs are marked by laCherry (magenta) and white arrowheads depict struts. The images are representative of 20 analysed embryos from two independent experiments. e, Chemical inhibition of the Notch signalling pathways by DAPT treatment or morpholino oligonucleotide (MO)-mediated knockdown of the downstream Notch target efnb2a in embryos in which arterial cells are marked by GFP (dll4) and all ECs by laCherry (magenta). As a control, a scrambled MO was injected at the same concentration as efnb2a MO or embryos were treated with the same concentration of DMSO in which the DAPT was dissolved. White arrowheads point at GFP (dll4) positive cells in the floor of the DA. Asterisks mark the disruption in arterial and venous unmixing. All images are representative of 20 analysed embryos from two independent experiments. f, A higher dose of efnb2a MO than in e results in a more dramatic unmixing phenotype. The magenta asterisk shows the disruption in arterial and venous unmixing and the red arrow shows the direction of blood flow. Note how some erythrocytes have faint laCherry expression, enabling visualization of the circulation. The images are representative of 30 analysed embryos from three independent experiments. Scale bars are defined in the figure.
Fig. 3 | Bone morphogenetic protein signalling is required for strut formation. a, Inhibition of BMP signalling by heatshock (hs) inducible expression of noggin (heatshock at 16 h.p.f.) or administration of 50 µM DMH1 from 16 h.p.f. onwards. The images are representative of 20 analysed embryos from two independent experiments. b, Ectopic expression of bmp2b by heatshock at 16 h.p.f. (upper) or 22 h.p.f. (lower). Asterisks indicate a partially formed CV wall, arrowheads indicate struts and arrows indicate a previously described venous sprouting phenotype\(^2\). The images are representative of 30 analysed embryos from three independent experiments. c, Stills of Supplementary Video 3: ectopic expression of bmp2b, heatshock at 16 h.p.f., prevents struts from pruning and consequently closure of the vessel wall. Arrowheads indicate struts and red and green arrows the migration direction of two ECs. The rightmost panel shows surface rendering with z-depth colour coding. The stills are representative of two independent time-lapse experiments in which one embryo is imaged each. d, Constitutively active Smad1 fused to mClover (green) under the control of the EC specific promoter fli1a was injected for mosaic expression in Tg(fli1a:laCherry) embryos in which all ECs are marked by laCherry (magenta). Arrowheads indicate the mClover-caSmad positive strut. The images are representative of 30 analysed embryos from three independent experiments. e, Different developmental stages of embryos in which all ECs are marked by laCherry (magenta). The images are representative of 20 analysed embryos from two independent experiments. Scale bars are defined in the figure.

the γ-secretase inhibitor DAPT\(^{14}\). In a complementary approach, we used a morpholino oligonucleotide (MO) to knock down the downstream Notch target ephrinB2a (efnb2a), which demarcates arterial and venous boundaries\(^{13-15}\). In both experiments, we found a dose-dependent inhibition of arterial–venous unmixing, resulting in a DA and PCV that remained fused (Fig. 2e and Extended Data Fig. 2a). The loss of Notch signalling in DAPT-treated embryos was illustrated by the marked reduction of dll4 expression, which we did not observe in efnb2a morphants, a downstream Notch target gene (Fig. 2e, arrowheads). Importantly, inhibition of arterial specification did not affect strut formation or pruning (Extended Data Fig. 2b). Thus, a single precursor vessel is formed by arterial and venous ECs in the CV region, with struts of either EC type, which over time unmixes into the DA and CV through increasing expression of arterial genes like efnb2a.

Because bone morphogenetic protein (BMP) plays a pivotal role in the remodelling of the CV into the CV plexus, a process that starts after the CV becomes functional\(^{14-20}\), we tested whether or not BMP signalling also plays a role in strut formation. To this end, we inhibited BMP signalling by expressing the BMP antagonist noggin via a Tg(hsp70:noggin) transgene\(^2\) or treatment with the small molecule DMH1, a dorsomorphin analogue\(^2\). Alternatively, we ectopically activated the bmp pathway by expressing bmp2b via the Tg(hsp70:bmp2b) transgene\(^2\). Noggin and bmp2b expression were controlled by the heatshock inducible promoter (hsp70) and thereby allowed for manipulation of BMP signalling after the BMP-dependent specification of the dorsoventral mesoderm\(^13,24\). Genetic or chemical inhibition of BMP signalling from 16 h.p.f. onwards resulted in a failure of the ECs to coalesce into struts (Fig. 3a). Next, we ectopically expressed bmp2b from 16 h.p.f. onwards and found that a portion of the struts were still present at 48 h.p.f. (Fig. 3b, top), whereas in wild-type embryos all struts were cleared from the lumen by 26–28 h.p.f. (Fig. 1b,c). In addition, we found that the CV wall in these embryos was only partially formed, which may be explained by the inability of ECs in struts to migrate into the CV wall (Fig. 3b, asterisk, 3c and Supplementary Video 3). Ectopic expression of bmp2b between 18 and 22 h.p.f. disrupts cardiac morphogenesis\(^2\) and, to circumvent this cardiac phenotype,
**Fig. 4** | Struts compartmentalize the caudal vein and trap erythrocytes. **a**, Visualization of erythrocytes by eGFP (*globin*) and all ECs by laCherry (magenta) in the trunk during strut formation. The images are representative of 30 analysed embryos from three independent experiments. **b**, Strut formation compartmentalizes the CV, thereby trapping erythrocytes into compartments. The images are representative of 30 analysed embryos from three independent experiments. **c**, Stills of Supplementary Video 4: pruning of struts results in the release of erythrocytes into the circulation. The stills are representative of five independent time-lapse experiments in which one embryo is imaged each. **d**, qPCR on dissected CV regions and FACS-sorted erythrocytes (10 CVs were pooled per condition measured in two independent experiments). **e**, qPCR on dissected CV regions from embryos injected with a scrambled MO or *tnnt2a* MO to prevent the onset of circulation (10 CVs were pooled per condition measured in two independent experiments). **f**, Embryos injected with a scrambled MO or *tnnt2a* MO. Arrowheads indicate struts that failed to prune. The images are representative of 20 embryos analysed from two independent experiments. The boxplot on the right shows the quantification of the number of struts at 48 h.p.f. (n = 15 for scrambled MO and n = 38 for *tnnt2a* MO injected; data are from two independent repeats and presented as the median values with the first and third quartiles; the whiskers represent the minimum and maximum. Two-tailed Student’s t-test. **P < 0.0001**). Scale bars are defined in the figure.
we heatshocked at around 22 h.p.f. Of note, at this time, the majority of the struts have already been pruned and therefore we could only target the few remaining struts (Fig. 1b,c). Similar to the experiments performed at 16 h.p.f., we also found struts that persisted at 48 h.p.f., while heart formation and circulation showed no obvious defects (Fig. 3b, lower). In addition, these embryos also displayed a previously described ectopic sprouting phenotype (Fig. 3b, arrows in bottom panel)19. Bmp2b regulates cellular responses through interaction with BMP receptors type I and II, resulting in the phosphorylation of Smads 1, 5 and 8 (R-Smads),
followed by heterodimerization of these phosphorylated R-Smads with co-Smad4 and ultimately the regulation of a myriad of downstream transcriptional targets, such as the angiogenic factor id1.25–27. To further test whether strut formation was facilitated through this canonical BMP pathway, we generated a constitutively active zebrafish Smad1 (caSmad1), placed under the control of the endothelial specific promoter fl1a, and injected it at the single cell stage for mosaic expression.28–30. Similar to the ectopic expression of bmp2b, caSmad1-mClover positive struts failed to pruned and were still present at 48 h.p.f., while injections of mClover alone did not prevent pruning of the struts (Fig. 3d and Extended Data Fig. 3a). By using a bmp reporter Tg(brecGFP), we found that besides expression in epidermal cells and erythrocytes, ECs in struts have active bmp signalling at different stages of CV formation (Fig. 3e). This is further supported by the presence of mRNA transcripts of both bmp2b receptors (bmpr2a and bmpr2b) and the downstream bmp target id1, indicative of active bmp signalling, in sorted ECs from the CV (Extended Data Fig. 3b). Thus, there is a temporal window in which bmp signalling is required for the formation of EC struts in the CV. Prolonging this window via ectopic activation of bmp signalling prevents struts from pruning.

Pruning of endothelial struts coincides with the onset of blood flow, an important factor in vascular remodelling and endothelial cell fate determination.25–28. To study blood flow in relation to endothelial struts, we simultaneously visualized erythrocytes and ECs during strut formation. At the onset of strut formation, erythrocytes were partially positioned between arterial and venous ECs and a large portion of the erythrocytes were mixed with venous ECs, especially in the CV region (Fig. 4a). As a result of this mixing, strut formation in the CV region trapped these erythrocytes into compartments, with some tightly packed with erythrocytes (Fig. 4b). The DA is initially devoid of erythrocytes (Fig. 4a,b; 20 h.p.f.) and they only start to enter the DA when blood flow is initiated (Fig. 4b; 24 h.p.f.). Although initially trapped in these compartments, erythrocytes are gradually released into the circulation when struts start to prune, resulting in progressive loss of these compartments (Fig. 4c and Supplementary Video 4). Erythrocytes have been shown to be a source of secreted growth factors and, given that they are closely associated with the ECs prior to strut formation, we questioned whether erythrocytes may be a source of the bmp2b that contributes to strut formation.29. To test this, we first used fluorescence-activated cell sorting (FACS) to sort erythrocytes at the time of CV formation and found bmp2b expression by quantitative polymerase chain reaction (qPCR) (Fig. 4d and Extended Data Fig. 4a). Of note, bmp2b is also expressed by the ventral mesoderm surrounding the CV, which is included in the unsorted population that we used as a reference.31. Next, we prevented erythrocytes from leaving the CV by inhibiting the onset of circulation by either blocking the expression of cardiac troponin T2a (tnnt2a) or through administration of the muscle relaxant ms-222 (tricaine methanesulfonate). From these embryos, we dissected the CV region and found bmp2b expression by quantitative PCR (Fig. 4e). We found no difference in the number of struts just before the onset of circulation (26 h.p.f.); however, at 48 h.p.f. we still observed the same number of struts as at 26 h.p.f. in embryos without circulation (Fig. 4f and Extended Data Fig. 4b). Furthermore, in these embryos the CV wall was also not fully formed, which is in line with our earlier findings that ECs from struts are reused to form the CV wall (Extended Data Fig. 4c, asterisks). In summary, we demonstrate that the erythrocytes secrete the Bmp2b that is required for strut formation, become trapped in compartments formed by struts and gradually enter circulation when these compartments are progressively lost through pruning of struts.

To test the functionality of struts, we ablated 1–2 cells within a single strut by ultrashort pulses of near-infrared laser light, which was sufficient to sever the strut. This technique has been shown to generate negligible heat transfer and collateral damage to neighbouring tissues.32,33. Because of the anatomical structure of the CV, we could only reliably sever endothelial struts after 22–24 h.p.f. (Fig. 5a and Supplementary Video 5). Although struts perpendicular to the sagittal plane were often difficult to detect and sever, we were able to ablate, on average, 95% of all struts within the CV (Fig. 5a, arrowhead in the middle right panel and Supplementary Video 6). As a control, we sham-treated embryos by positioning the laser 25–50 μm from struts, in the empty space of the lumen, applying the same number of ablation pulses as in treated embryos. Severing of nearly all endothelial struts initially did not lead to a dramatic change in the shape of the lumen. However, at the onset of circulation, the lumen of the CV collapsed due to a nearly instantaneous removal of the erythrocytes from the CV (Fig. 5b, Extended Data Fig. 5a and Supplementary Video 7). This collapsed CV is severely mispatterned during the remodelling of the CV into the CV plexus, although it allowed some blood flow (Fig. 5c). In animals in which circulation has just initiated, ablating a single strut often resulted in the release of erythrocytes into the circulation, demonstrating that these compartments trap erythrocytes (Supplementary Video 5, embryo 2).

In summary, we have shown that strut formation traps erythrocytes into compartments, with some tightly packed with erythrocytes, and thereby forms a rigid network that withstands external forces and maintains the shape of the vessel. Pruning of struts occurs gradually, ensuring that not all erythrocytes enter the circulation simultaneously, which would result in collapse of the vessel (Extended Data Fig. 5b).

Online content

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Methods

Zebrafish husbandry. Zebrafish (Danio rerio) were maintained and propagated according to the guidelines of the UCSD Institutional Animal Care and Use Committee and Huberecht Institute Ethical Review Board. All animal experiments were approved by the Animal Experimentation Committee (DEC) of the Royal Netherlands Academy of Arts and Sciences. Embryos and adult fish were raised in a circulating aquatic system (Aquaneering) at 28 °C. The following zebrafish lines have been described previously: Tg(BIhlfluorescentCherry)772 (ref. 37); referred to as fltlaaCherry; Tg(fltaaGFP)83 (ref. 38); Tg(hb1.1GFP)88 (ref. 39); referred to as Tg(fltdolecGFP); Tg(RLkDENDRA2) (ref. 40); Tg(BLkGaiAEP)40 (ref. 41); Tg(UAS:fluorescentGFP)255 (ref. 42); referred to as UAS:GFP; Tg(EPVTp1:mNnu). HtbGFP:stra3 (ref. 43) referred to as Tg(Tp1:GFP); Tg(hsp70bmp2b)186 (ref. 44); Tg(adrlinGFP)40 (ref. 45). Details on the strains and ages are noted for each experiment. Gender was not selected in any of the studies conducted.

Morpholino and plasmid injections, heat shock and chemical treatment. Embryos were injected at the one-cell stage with MOs (GeneTools) or 100 ng of plasmid with 100 ng of tol2 mRNA. EphrinB2a (efn2a) translation blocking MO (5'-CGGCTAATACCTTGTGTCGCGGA-3') silences heart morphogen trout2a(5'-CATGTGTCGACCTACGTGGAC-3') expression. Scrambled MO was injected at similar concentrations as the targeted MO (5'-CCCTCTTACTGCTATTACAACTAAATA-3'). Capped tol2 mRNA was synthesized from linearized pCS2 constructs using the mMessage mMachine ′机器′ (Ambion, AM1340). Injected and un-injected control embryos were heatshocked at 38 °C for 30 min. Embryos were treated with 50 μM DMH1 (#166797, Cayman Chemical; DAPT (all-Cayman Chemical) dissolved in DMSO (1.000x stock solution) and control embryos were treated with DMSO alone. Blood flow was inhibited by treating the embryos with 0.48 mg ml⁻¹ (3x) ethyl 3-aminobenzoate methanesulfonate (MS222; Sigma E10521) at a temperature of 28.5 °C. Imaging was performed using SP5 (Leica) or a Zeiss LSM 880 with an Airyscan system. For all imaging, live microscopy was carried out using a confocal microscope or a Zeiss LSM 880 with an Airyscan system. For all imaging, live microscopy was carried out using a confocal microscope or a Zeiss LSM 880 with an Airyscan system. For all imaging, live microscopy was carried out using a confocal microscope. Scoring was performed using high-resolution longitudinal imaging of the tail and posterior trunk of larval zebrafish. Zebrafish 14, 489–491 (2017).

Cloning. The coding sequence of zebrafish Small1 (ENSARG00000027199) was cloned into the pcS2+ vector by the Gibson cloning method (NEB HiFi DNA assembly mix, E0521). To substitute serine 470 and 472 into aspartic acid (SVS to DVS), zsMad1 was cloned with reversed primers containing the required Nucleotide changes (see Supplementary Table 1 for the primer sequences) and fused to mClover3 (N-terminally) flexibly linked to GSAAGAAGAGEF in front of the fltlaa endothelial specific promoter, referred to as Tg(flt1:GFP)144 (ref. 46). The coding sequence of EphrinB2a (ENSARG00000020164) was cloned into the pcS2+ vector and mRNA was transcribed with the Sp6 message machine kit (Promega).

Cell preparation and flow cytometry. Embryos were collected and anaeusthetized in E3 medium containing 0.01% tricaine, then the CV region or trunk region was dissected and placed in ice-cold PBS. After centrifugation, the pellet was washed with ice-cold PBS and resuspended in TrypLE (Thermo Fisher, 1253011) for 45 min at 32 °C with vigorous shaking. Cells were then washed with FACS buffer (PBS + 2% FBS + 2 mM EDTA), dissolved in FACS buffer + DAPI (Thermo Fisher, 0.5μg/ml final concentration) and subsequently filtered through 40-μm nylon mesh. Cells were sorted on a FACSAria II system (Becton Dickinson) based on bright fluorescence collected in FACS buffer for gene expression analysis. Forward scatter (FSC) and side scatter (SSC) were cut at 50k and live cells were DAPI negative (<5x10⁶). The gating strategies for erythrocytes (GFP) and endothelial cells (mCherry) are shown in Extended Data Figs. 3b and 4a.

Quantitative PCR and PCR with reverse transcription. RNA was extracted from dissected tissue (CV region) or sorted cells using Qiazol (Thermo Fisher), according to the manufacturer’s instructions. cDNA was generated using GoScript (Promega) using random hexamer primers. qPCR assays were performed on a Bio-Rad CFX96 real-time system according to the manufacturer’s instructions (Bio-Rad) with Fast SYBR Green (Bio-Rad). The expressions of ef1a, 18s, btp and bactin were used to normalize the amount of the investigated transcripts. The primers used are listed in Supplementary Table 1.

Imaging. Images captured in Arysyan mode (Zeiss LSM 880 with Zeiss ZEN blue version 3.0) were first processed in Zeiss ZEN blue (Arysyan processing) and subsequently analysed using Fiji (version 1.52p) or Imaris (version 9.1). Sub-stacks were subtracted to specifically visualize the blood vessel wall or lumen, or z-stacks were flattened by the maximum intensity projection method used as stated in the figure captions. In some cases, XYZ slices were corrected using the MultiStackReg plugin (Fiji; B. Busse, NICHD, version (1.15)) and correction of fluorescence bleaching by histogram matching (Fiji). Tiled images were stitched with either Zeiss ZEN or Imaris sticher (version 9.1). Contrast in all images was adjusted and look-up tables (LUT) adjusted (red to magenta or red/green to black and white) in Fiji or Imaris for visualization purposes. Surface renderings were performed in Imaris. Figures, videos and animations were created with Adobe Illustrator, Photoshop, After Effects and Premier (version CS5).

Reagents. A list of reagents and catalogue numbers is provided in Supplementary Table 2.

Statistical analysis and experimental set-up. For each experiment, animals from the same clutch were divided into different treatment groups without any bias. The whole clutch was excluded if more than 10% of control embryos displayed obvious developmental defects. If necessary, statistical analysis was performed using SPSS version 20 (IBM). The Mann–Whitney U test was used for statistical analysis of two groups with unequal variances. The unpaired t-test was used for two groups with equal variances. The Kruskal–Wallis test was used for statistical analysis of multiple groups with equal variances, and one-way analysis of variance for multiple groups with unequal variances. Dunn’s post-hoc test was used for pairwise multiple comparisons. All experiments are a representation of at least two independent repeats.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this Article.

Data availability

Source data are provided with this paper. All other data supporting the findings of this study are available from the corresponding author upon reasonable request.

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Author contributions
B.W. designed, performed, analysed experiments and wrote the manuscript. Laser ablation experiments were performed by B.W. and I.S. under the supervision of D.K. M.G. interpreted data and designed experiments. Part of the project was supervised by C.R. D.T. supervised the project and wrote the manuscript. All authors discussed the results and commented on the manuscript at all stages.

Competing interests
The authors declare no competing interests.

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Correspondence and requests for materials should be addressed to B.W. or D.T.
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Extended Data Fig. 1 | Characterization of endothelial struts. a, All ECs are marked by both laCherry (magenta; cytoskeleton) and eGFP (green; nuclei). Struts are depicted by arrowheads. Images are representative of 20 embryos analysed from two independent experiments. b, Quantification of the number of endothelial struts at 22 hpf and 26 hpf by confocal microscopy and represented as the median values with the first and third quartiles; the whiskers represent the minimum and maximum (4 different clutches per time point (n = 26; n = 25; n = 27; n = 25 for 22 hpf and n = 25; n = 28; n = 25; n = 25 for 26 hpf. Two-tailed Student’s t-test. ***P < 0.0001 for all 4 tests). Arrowheads depict endothelial struts. Images are representative of indicated groups from four independent experiments. c, Strut formation in the anterior part of the PCV at 22hpf by confocal microscopy and represented as the median values with the first and third quartiles; the whiskers represent the minimum and maximum (n = 30 embryos from three independent experiments). Arrowheads depict endothelial struts. d, Quantification of the number of acridine orange positive ECs in the CV region at 20 hpf and representative confocal images. White arrowhead depicts a positive EC (n = 20 embryos from two independent experiments). Scale bars are defined in the figure.
Extended Data Fig. 2 | Unmixing of arterial and venous endothelial cells. a, Rescue of ephrinb2a morphants with efnb2a mRNA. Embryos were grouped by the level of rescue (near full, partial or no rescue) of the floor of the DA, indicated by the asterisks. (n = 71 embryos analysed in one experiment, n numbers per group indicated in graph). Images are representative of 5 embryos analysed by confocal microscopy per group. b, Representative images and quantification of the number of struts in embryos treated with the Notch inhibitor DAPT or injected with an ephnb2a MO. As a control, embryos were injected with a scrambled MO. (n = 14 embryos per condition). Data presented as the median values with the first and third quartiles; the whiskers represent the minimum and maximum. Images are representative of 5 embryos analysed by confocal microscopy per condition. Scale bars are defined in the figure.
Extended Data Fig. 3 | BMP signalling is required for strut formation. a, Mosaic expression of mClover under the control of the EC specific promoter fli1a. Arrowhead indicates mClover positive strut at 22 hpf (upper panel). Lower panel shows an image of the same region at 48 hpf. Images are representative of 9 embryos analysed from three independent experiments. b, Gating strategy used to FACS sort ECs from dissected CV regions followed by qPCR (10 CVs were pooled per condition measured in two independent experiments). Of note, the bmp pathway is active in multiple tissues at this time, including erythrocytes and strong activity in epithelial cells (Fig. 3e asterisks), these cells are included in the unsorted CV (black bars). Scale bars are defined in the figure.
Extended Data Fig. 4 | Inhibition of blood flow prevents struts from pruning. a, Gating strategy used to FACS sort ECs from dissected CV regions followed by qPCR (b). The onset of blood flow was inhibited by either injecting of a cardiac troponin T2a (tnnt2a) MO or through administration of 3x concentrated ms-222 (tricaine methanesulfonate). Erythrocytes are marked by GFP (globin) and all ECs by laCherry (magenta). Data are presented as the median values with the first and third quartiles; the whiskers represent the minimum and maximum. One-way analysis of variance (ANOVA) with Dunn's post-hoc test. ns = not significant (n = 25 embryos per time points per experiment of three independent experiments). Images are representative of 5 embryos analysed by confocal microscopy per condition per experiment. c, 48 hpf embryo injected with tnnt2a MO. Asterisks indicate an incomplete formation of the CV wall. Images are representative of 10 embryos analysed from two independent experiments. Scale bars are defined in the figure.
Extended Data Fig. 5 | Endothelial strut model. a, Laser ablation of a single strut resulted in a slight deformation of the CV, shown by the superimposed image of all timepoints. Stills are representative of 18 embryos analysed from three independent experiments. Scale bar is defined in the figure (b). Schematic. 1) Venous (blue) and arterial (red) EC precursors originate from distinct locations within the lateral plate mesoderm, with the primitive erythrocytes (precursors) positioned medially adjacent to the venous ECs. 2) ECs and erythrocytes migrate towards the midline of the embryo. Arterial ECs migrate along and directly contact the ventral face of the somites, thereby they receive inductive cues that strengthens the arterial fate (shades of red). 3) At the midline, venous and arterial ECs coalesce into a network of struts and form a common precursor vessel. This process is bmp2b dependent, which is secreted by, among others, the erythrocytes. Strut formation encloses erythrocytes into compartments. 4) The vessel wall consists initially only out of a few patches of ECs and upon pruning of struts, ECs from struts migrate into and are incorporated in the vessel wall. 5) Arterial ECs participating in strut formation have a weak arterial identity, which progressively increases (shades of red) and results in the expression of the Notch target efnb2b, which drives the unmixing of arterial and venous ECs. Segregation of these arterial and venous ECs results in the formation of the DA and CV. 6) Onset of circulation flushes the bmp2b expressing erythrocytes from the CV, which is an important step for complete pruning of all struts. The lumen of the CV is now maintained by blood pressure rather than through the support of struts.
Reporting Summary

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Our web collection on Statistics for Biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection
- Zeiss Zen blue (version 3.0)
- Leica LAS X (version 3.4.2.18368)

Data analysis
- Zeiss blue (version 3.0)
- Leica LAS X (version 3.4.2.18368)
- Fiji (version 1.52p)
- Fiji plugin MultiStackReg (to correct for drift version 1.45)
- Imaris (version 9.1)
- Imaris Sticher (version 9.1)
- R Statistical Package Version 3.4.0, R Core Team
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All data supporting the findings of this study are available from the corresponding author on reasonable request.

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size
Sample size (approximately n=20) were chosen based to garner sufficient confidence between groups (Mead’s Resource Equation). Animals were randomly picked from the entire clutch.

Data exclusions
No data was excluded at the post-experimental stage. At the pre-experimental stage, a clutch was excluded if more than 10% of the embryos displayed obvious developmental defects.

Replication
All experiments were repeated at least twice (different day and breeding pair). All imaging data were successfully repeated by using different reporter lines (e.g. flI1a:GFP, flI1a:CherryCAAX, flI1a:LifeactCherry, KDR:eGFP). Except if the was no alternative transgenic line available: photoconversion (flk1:DENDRA, Fig. 1d and e), Notch reporter (TP1:eGFP, Fig. 2d), BMP reporter (8RE:eGFP, Fig. 3e)

Randomization
For each in vivo experiments, animals from the same clutch were divided into different treatment group without any bias (randomly distributed)

Blinding
Blinding was not performed as treated animals (e.g. chemically with BMP inhibitor or heat shock with noggin) can be easily distinguished from their control clutch mates due to their strong phenotype

Reporting for specific materials, systems and methods

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Methods

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| ☒   | ChiP-seq |
| ☒   | Flow cytometry |
| ☒   | MRI-based neuroimaging |

Animals and other organisms

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Laboratory animals

Zebrafish (Danio rerio) were maintained and propagated according to the guidelines of the UCSD Institutional Animal Care and Use Committee (IACUC) and Hubrecht Institute ethical review board. All animal experiments were approved by the Animal Experimentation Committee (DEC) of the Royal Netherlands Academy of Arts and Sciences. Embryos and adult fish were raised in a circulating aquarium system (Aquaneering) at 28°C. The following zebrafish lines have been previously described:

Tg[flI1a:LifeactCherry]vnc7Tg [ref 18] referred to as flI1a:Cherry; Tg[flI1a:GFP]y1 [ref 30]; Tg[hba:1.1:EGFP]446 [ref 37] referred to as Tg[hba:1.1:EGFP]; Tg[flk:DENDRA2] [ref 38]; Tg[dl4:G.a:4.FP]hu10049Tg [ref 12]; Tg[UAS:LifeactGFP]mu271 [ref 35] referred to as
Flow Cytometry

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Methodology

Sample preparation

Embryos were collected and anesthetized in E3 medium containing 0.01% tricaine the CV region or trunk region was dissected and placed in ice-cold PBS. After centrifugation the pellet was washed with ice-cold PBS and resuspended TrypLE (ThermoFisher, 12563011) for 45min at 32 °C with vigorous shaking. Cells were then washed with FACS buffer (PBS + 2% FBS + 2mM EDTA), dissolved in FACS buffer + DAPI (ThermoFisher, 0.5ug/ml final concentration) and subsequently filtered through 40-μm nylon mesh. Cells were sorted on a FACS Aria II (BD Biosciences) and collected in FACS buffer for gene expression analysis.

Instrument

FACS Aria II (BD Biosciences)

Software

BD FACSDiva (version 8.0)

Cell population abundance

A minimum of 10,000 and maximum of 15,000 cells were sorted for mRNA extraction followed by quantitative PCR

Gating strategy

FSC and SSC were cut at 50K, live cells were DAPI negative (< 5x10^4%). Gating strategy for erythrocytes (GFP) and endothelial cells (mCherry) are shown in Extended Data Fig. 3b and 4a

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