Single-Cell Transcriptional Profiling of Aortic Endothelium Identifies a Hierarchy from Endovascular Progenitors to Differentiated Cells

Graphical Abstract

Highlights

- scRNA sequencing reveals two distinct aortic endothelial populations
- Sequencing analysis confirms transitional cells rather than discrete populations
- Endovascular progenitors are slow cycling and have distinct mitochondrial content

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In Brief

Lukowski et al. demonstrate the existence of two distinct endothelial cell populations in the aorta via single-cell sequencing. The data confirm that a progenitor population transitions to a mature endothelial cell, defining an endothelial hierarchy.
Single-Cell Transcriptional Profiling of Aortic Endothelium Identifies a Hierarchy from Endovascular Progenitors to Differentiated Cells

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SUMMARY

The cellular and molecular profiles that govern the endothelial heterogeneity of the circulatory system have yet to be elucidated. Using a data-driven approach to study the endothelial compartment via single-cell RNA sequencing, we characterized cell subpopulations within and assigned them to a defined endothelial hierarchy. We show that two transcriptionally distinct endothelial populations exist within the aorta and, using two independent trajectory analysis methods, confirm that they represent transitioning cells rather than discrete cell types. Gene co-expression analysis revealed crucial regulatory networks underlying each population, including significant metabolic gene networks in progenitor cells. Using mitochondrial activity assays and phenotyping, we confirm that endovascular progenitors display higher mitochondrial content compared to differentiated endothelial cells. The identities of these populations were further validated against bulk RNA sequencing (RNA-seq) data obtained from normal and tumor-derived vasculature. Our findings validate the heterogeneity of the aortic endothelium and previously suggested hierarchy between progenitor and differentiated cells.

INTRODUCTION

The circulatory system throughout an organism is lined by specialized endothelial cells that form the lumen of the vasculature tube (Herbert and Stainier, 2011). It is becoming apparent that, within each organ bed, the endothelial compartment is heterogeneous. This includes variations between vessel types (arterial, venous, and lymphatic), location (macro- or microvascular), and between neighboring endothelial cells (Adams and Alitalo, 2007; Eichmann et al., 2005), with multiple subpopulations being found, such as tip and stalk cells (Iruela-Arispe and Davis, 2009). Although each of these situations has been examined and validated individually on a candidate-based approach, there has never been an unbiased attempt to describe the heterogeneity that characterizes this compartment (Aird, 2012). Furthermore, although each aspect of heterogeneity has been explored individually, the overlap between the various indicators of heterogeneity is currently unknown. Importantly, the complexity in vessel regeneration or turnover compounds the need to better understand the interactions between the endothelial populations and ascertain the specific genes that control these processes (Aird, 2007a, 2007b).

We recently reported that in a variety of vascular beds reside an endothelial stem cell population that was termed endovascular progenitor (EVP) (Patel et al., 2017). These cells are governed by a stringent set of functional characteristics, such as slow cycling, self-renewal, engraftment potential, and finally differentiation to transit amplifying and mature endothelial cells. In the initial description, making use of common endothelial markers, these EVP cells had a stringent cell surface marker expression profile, CD34+CD45−CD31loVEGFR2−, and also gave rise to a transit amplifying (TA) population, which then transitioned to a definitive differentiated (D) endothelial cell with increasing surface levels of VEGFR2 and CD31. This endothelial hierarchy was observed not only in situations of homeostasis (aorta) but also of pathology, such as wound healing or cancer. Bulk RNA sequencing was conducted on EVP and D cells from the aorta, demonstrating remarkable gene expression changes during the transition. However, others have described different progenitor populations with endothelial potential characterized by the expression of different genes, such as CD157, c-Kit, or activated protein C (Fang et al., 2012; Wakabayashi et al., 2018; Yu et al., 2016). It is therefore difficult to establish the relationship between these different cell populations emanating from different studies.
The recent development of single-cell RNA sequencing (scRNA-seq), which combines single-cell isolation techniques with RNA-seq, creates an opportunity to study the transcriptomes of individual isolated cells. This enables clear distinction to be made between subpopulations and the thorough assessment and quantification of gene transcripts in an unbiased way without preconception. In this study, we aimed at characterizing the gene expression in the endothelium of the aorta at single-cell resolution to obtain a clear and unbiased picture of endothelial cell populations. Our findings largely confirm the previously described hierarchy between a slow-cycling EVP and a terminally differentiated D population, opening an understanding of the homeostasis of the vasculature.

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**RESULTS**

### scRNA-Seq

We obtained an unbiased study of the aorta endothelium by dissecting the descending aorta of 6- to 8-week-old mice and proceeded to cell separation of the endothelium, based on living Lin+CD34+ cells (Figures 1A and 1B). We further demonstrate that >99% of this population of cells express VE-cadherin (VECAD) to highlight their restricted endothelial lineage (Figure 1C). Sequence data from three scRNA-seq libraries constructed from three pools of 30 mouse aorta samples were combined for processing and analysis. Pre-quality control (QC) libraries contained 2,212, 3,256, and 2,203 cells, a total of 7,671 cells. We detected a median of 1,850 unique molecular identifiers (UMIs) per cell, a median of 788 genes per cell, and obtained an average read depth of 53,631 reads per cell. This sequencing depth allowed us to reach saturation for the number of detected genes per cell in each of the 3 samples (Figure S1). Following quality control processing in the ascend package, we obtained 1,638, 2,821, and 1,652 cells for a total of 6,111 high-quality cells in the final dataset.

### Identification of Distinct Endothelial Clusters in Aorta Endothelium

Unsupervised hierarchical clustering analysis of the scRNA-seq data using the Clustering at Optimal Resolution (CORE) method (Nguyen et al., 2018) revealed four stable cell populations containing 3,231 (52.87%), 2,636 (43.14%), 164 (2.68%), and 80 (1.31%) cells, respectively. These clusters were stable across
pressed genes per cluster sorted by mean expression and log2 as a contaminant. Table 1 shows the top differentially expressed genes, such as Sox17, Cdh5, Fabbp4, and Cldn5 (Figure 1F). On the other hand, cells in cluster two displayed a more mesenchymal phenotype and overexpressed genes, such as Dcn, Lum, and Gsn (Patel et al., 2017). Cells belonging to cluster three also displayed a distinct terminally differentiated endothelial phenotype but also expressed markers of inflammation, including Il1b and Atf1, compared to other endothelial cells in clusters one and two. Cells in cluster four were identified as either cardiac or skeletal muscle due to their high expression of troponin and other myocyte markers, including Tnn2, Tnni3, and Myf7 (cardiac) and Tnrc2, Tnn3, Myf1, and Myf4 (striated muscle). Although of interest, this fourth cluster represented only a small fraction of the entire sample, and its relevance to endothelial biology was uncertain. Hence, we proceeded to further analysis by considering this fourth cluster as a contaminant.

Table 1. Top 20 Differentially Expressed Genes per Cluster (Cluster x versus Remaining Clusters) Sorted by Highest Mean Expression and Log2 Fold Change

| C1 Gene | C1 log2FC | C2 Gene | C2 log2FC | C3 Gene | C3 log2FC | C4 Gene | C4 log2FC |
|---------|-----------|---------|-----------|---------|-----------|---------|-----------|
| Jchain  | 6.61      | Serpinf1| 2.78      | H2-Ab1  | 6.16      | Tnnc2   | 10.33     |
| Cldn5   | 3.85      | Lum     | 2.78      | Hmgb2   | 6.08      | Acta1   | 10.28     |
| Rgs5    | 3.76      | Mmap5   | 2.62      | H2-Ab1  | 5.89      | Slr      | 10.24     |
| Egf7    | 3.73      | Smoc2   | 2.59      | H2-Aa   | 5.87      | Myf7    | 10.15     |
| Fabbp4  | 3.66      | Pcolce  | 2.54      | Cdt7    | 5.36      | Myf1    | 9.98      |
| Ccl5    | 3.58      | Gsn     | 2.47      | H2afz   | 4.86      | Myf4    | 9.88      |
| Rbp7    | 3.57      | Ccl11   | 2.47      | Hmgb1   | 3.43      | Tntt2   | 9.82      |
| Gpibbp1 | 3.55      | Dcn     | 2.47      | Ptau    | 3.38      | Ckm     | 9.81      |
| Esam    | 3.31      | Clc3b   | 2.46      | Tmsb4x  | 3.00      | Mb      | 9.73      |
| Ccld4   | 3.25      | Nupr1   | 2.36      | Cst3    | 2.93      | Tnnt3   | 9.73      |
| Hbb-bt  | 3.03      | Pf16    | 2.36      | Pf1     | 2.67      | Myf3    | 9.18      |
| Cd36    | 2.93      | Col1a2  | 2.36      | Actb    | 2.80      | Actc1   | 9.18      |
| Hba-a2  | 2.88      | Col3a1  | 2.30      | Actg1   | 2.68      | Fabp3   | 9.13      |
| Hba-a1  | 2.85      | Serping1| 2.29      | Ppia    | 2.62      | Cx6a2   | 8.86      |
| Hbb-bs  | 2.58      | Igfbp6  | 2.23      | Gm9843  | 2.47      | Tpm1    | 8.16      |
| Rgcc    | 2.37      | Tmp2    | 2.17      | Fau     | 2.38      | Slc25a4 | 5.22      |
| Tspan13 | 2.33      | Rarres2 | 2.14      | Crip1   | 2.20      | Atg5g1  | 4.35      |
| Id1     | 2.26      | Serpinh1| 2.13      | Etf1a1  | 2.16      | Atg5e   | 3.04      |
| Aap1    | 2.21      | Cdh63   | 2.10      | Tmsb10  | 2.08      | Cx7c    | 2.87      |
| Cyt1    | 2.17      | Gpx3    | 2.01      | Oaz1    | 2.00      | Cx6c    | 2.84      |

Expression Differences between EVP and D Populations

Previously, we detected EVP progenitors and D differentiated cells in aortic endothelium and tumor tissue using whole-tissue RNA-seq (Donovan et al., 2019; Patel et al., 2017). These populations were shown to express a set of genes that place cells along the continuum of differentiation (Sox9, Il33, and Pdgrfa in EVP and Pecam1 and Sox18 in D). To further investigate the transcriptional differences between populations at single-cell resolution, we restricted our analysis to clusters one and two. A differential expression analysis between these subpopulations revealed 234 genes (log2 fold change > 2; adj. p value < 0.01; Figure 1D). Of these, 90 were upregulated in cluster one and 144 were upregulated in cluster two. We observed cells in cluster one expressed Pecam1 (log2 fold change = 3.06; adj. p = 2.53 × 10−68) and Cdhs5 (log2 fold change = 2.91; adj. p = 4.05 × 10−14) at significantly higher levels compared to cells in cluster two as observed in the D population. Conversely, cells in cluster two showed increased expression of Pdgrfa (log2 fold change = 2.00; adj. p = 1.02 × 10−85), Il33 (log2 fold change = 2.06; adj. p = 3.46 × 10−15), and Sox9 (log2 fold change = 2.18; adj. p = 8.94 × 10−7), as observed in EVP (Patel et al., 2017).

Using these marker genes, we visualized cells belonging to clusters one and two using t-Distributed Stochastic Neighbor (t-SNE) dimensionality reduction and observed a clear separation between cells expressing Cdhs5 and Pecam1 (D; cluster one) and those expressing Sox9, Il33, and Pdgrfa (EVP; cluster two; Figure 2A). These findings indicate that clusters one and two correspond to transcriptionally distinct populations of endothelial cells at different stages of differentiation.

Marker genes derived from the literature are used to establish cell identity and infer function of clustered cells; however, many canonical marker genes do not clearly show subtle changes in the three batch experiments, giving confidence that they represented the biology of the homeostatic aortic endothelium (Figures 1D and 1E). To identify the cell types represented by each cluster, we performed differential gene expression analysis, comparing the cells in each cluster against the remaining cells. Our results revealed that cells in cluster one displayed a mature, differentiated vascular endothelial signature with increased expression of specific genes, including Sox17, Cd36, Fabbp4, and Cldn5 (Figure 1F). On the other hand, cells in cluster two displayed a more mesenchymal phenotype but also expressed markers regulated in cluster two. We observed cells in cluster one expressed Sox9 (log2 fold change = 2.18; adj. p = 8.94 × 10−7).
gene expression that exist between closely related cell populations. Machine learning approaches have been used to refine gene lists that capture the transcriptional signature of cell populations in scRNA-seq analyses in order to produce a list of genes that explain the greatest amount of deviance (Nguyen et al., 2018). Using a least absolute shrinkage and selection operator (LASSO) regression approach, we further characterized each cell cluster and established a set of gene markers describing the transcriptional differences between each of the four clusters. Starting with differentially expressed genes obtained for each individual cluster, we used the LASSO regression procedure to detect gene markers for each of the four clusters, obtaining 162, 135, 33, and 25 genes for clusters 1–4, respectively. Our results show that each cluster expresses a subset of genes

Figure 2. Expression of Marker Genes Defining Each Population

(A) Expression levels of six canonical marker genes were overlaid on the 2D t-SNE plot and highlight the cell specificity of their expression. The expression levels are shown as log_{2}(counts + 1)-transformed values, and the gradient represents low (gray) to high values (purple).

(B) Heatmap of LASSO-identified marker genes. Marker genes identified using the LASSO machine learning approach clearly mark each cluster and explain the most deviance per cluster. Expression levels are plotted as log_{2}(counts + 1)-transformed values. The color gradient shows low (purple) to high expression levels (yellow).

(C) Pairwise Spearman’s correlations were performed between single-cell data for each cluster and bulk RNA-sequencing data for normal or tumor populations of D and EVP. The correlation coefficient for each test is shown in the center of each square, and the color gradient shows the range of correlation coefficients from 0 (blue) to 1 (red).
distinguishing it from the remaining clusters. For example, cluster one (D cells) expressed higher levels of fatty acid binding protein 4 (Fabp4) and Jchain, and cluster two (EVP) showed increased expression of several serpins (Serpinf1, Serpinf1, and Serpinh1), Cct11, Postn, and Dcn. Cluster three had markedly higher expression of important transcription or translation factors, including H2az2, Bf13, Hif3alpha, and Eefl1a1, as well as Ppia, which encodes cyclophilin A. Finally, cluster four cells expressed genes related to cardiac muscle, including Mb, Tnn3, and Tnn1c1, which confirmed our initial findings. Expression levels of across all clusters are shown for 103 genes in Figure 2B, and the table of genes per cluster and their associated deviance coefficients are presented in Table S1.

Correlation of Whole-Tissue D and EVP to scRNA-Seq Clusters One and Two
The single-cell transcriptional profiles that we generated allow a high-resolution investigation of endothelial subpopulations. Our previous study of EVP and D populations in aortic endothelium used a bulk RNA-seq approach to examine gene expression in whole-tissue samples (Patel et al., 2017). To confirm that the cell populations in single-cell clusters one and two were consistent with the whole-tissue data D and EVP, respectively, we performed pairwise Spearman’s correlations between single-cell clusters and bulk transcriptome data. The gene expression profiles of each were compared using the significantly differentially expressed genes marking each of the four major scRNA-seq populations or the whole-tissue EVP and D populations, a total of 2,105 genes. We observed a very strong correlation for cluster two (proposed EVP population) with the whole-tissue EVP (Spearman’s rho = 0.83; p < 2.2 × 10^{-16}), and the cluster one (proposed D population) cells were well correlated with whole tissue D (Spearman’s rho = 0.64; p = 1.28 × 10^{-239}). We also observed a moderate, albeit less significant, correlation of whole-tissue EVP with cluster one cells (Spearman’s rho = 0.60; p = 9.03 × 10^{-202}, Figure 2C).

We also tested bulk RNA-seq data that we generated for EVP and D endothelial populations isolated from a B16 model of melanoma (Donovan et al., 2019). In these tumor vasculature populations, we also observed a strong correlation of tumor EVP with normal tissue EVP (Spearman’s rho = 0.75; p < 2.2 × 10^{-16}) and cluster two (Spearman’s rho = 0.78; p < 2.2 × 10^{-16}), and the tumor D population correlated well with normal tissue D (Spearman’s rho = 0.61; p < 2.2 × 10^{-16}) and cluster one (Spearman’s rho = 0.58; p = 4.28 × 10^{-186}). We also noted the tumor EVP population correlated with cluster one and may reflect higher heterogeneity and/or deranged gene expression in the tumor. Of interest, cluster three best correlated with tumor D populations, suggesting that it consists of a population of differentiated cells (D) that can be observed in situations of injury.

Together, the results of the pairwise correlation analysis further support the classification of cluster one as the D endothelial cell population and cluster two as the EVP population.

Linear Modeling of Gene Expression in D versus EVP Populations
To identify further biologically important differences between the D and EVP populations, we applied a linear modeling approach to investigate changes in the expression levels of each non-zero expressed gene (n = 17,175) per individual cell (m = 5,867) against their assigned cluster, either cluster one (D) or cluster two (EVP). A Bonferroni correction for multiple testing was used to determine statistical significance per gene (p = 2.89 × 10^{-15}), and from these, we retained 4,616 significant results. Our data revealed the top genes with significant changes in expression between D and EVP populations, including Gsn (beta = 52.5; p < 2.2 × 10^{-15}), Dcn (beta = 25.6; p < 2.2 × 10^{-15}), S100a6 (beta = 10.6; p < 2.2 × 10^{-15}), Fth1 (beta = 10.1; p < 2.2 × 10^{-15}), Fth1 (beta = 11.0; p < 2.2 × 10^{-15}), Fabp4 (beta = -11.8; p = 3.7 × 10^{-15}), Hbb-bs (beta = -37.1; p = 2.4 × 10^{-6}), Hba-a1 (beta = -27.3; p = 6.1 × 10^{-7}), and Hba-a2 (beta = -23.7; p = 7.3 × 10^{-5}). Overall, we detected 4,167 genes with significantly increased expression in EVP cells (mean beta = 0.23) and 449 with increased expression in D cells (mean beta = -0.34). We noted that several of these genes were also detected as innovative marker genes using the aforementioned LASSO method. Here, a positive beta value indicates that, for each unit change in gene expression in D, expression was increased in EVP by the beta value and vice versa. The results of the linear modeling analysis are presented in Table S1.

Co-expressed genes in the D or EVP populations are likely to form regulatory networks specific to the cell type. Weighted gene correlation network analysis (WGCNA) (Langfelder and Horvath, 2008) was performed using all 449 upregulated D genes and the top 449 (out of 4,167 total) EVP genes, and two robust modules were detected in each cell population (Figures 3A and 3B). The D modules contained 194 (blue) and 139 (orange) genes (Figure 3A), and EVP modules contained 121 (blue) and 257 (orange; Figure 3B). We tested whether the genes in each module share important biological functions and observed significant Gene Ontology (GO) term enrichment for cell-type-specific biological processes. In D cells, we identified significant genes underlying endothelial cell differentiation and vasculogenesis (Figures 3C and 3E). The genes in the EVP modules were enriched for cell metabolism, extracellular matrix organization, and epithelial cell proliferation (Figures 3D and 3F). StringDB (Szklarczyk et al., 2015) analysis further confirmed the enrichment of functions tightly linked to cell adhesion, cell differentiation, and endothelium development (D) and metabolism and vasculature development (EVP; Figures S2, S3, S4, and S5).

Using the top 449 genes from the linear regression analysis with negative (D) or positive (EVP) beta values, we interrogated the Enrichr database (Chen et al., 2013) to determine whether those genes increased in D or in EVP represented known cell or tissue types. For genes upregulated in D cells, we observed a highly significant overlap with human umbilical vein endothelial cells (HUVECs) (Z-score = -1.55; adj. p = 4 × 10^{-41}), and for genes upregulated in EVP cells, we observed a similar but less significant overlap reflecting their endothelial origin (Z-score = -1.44; adj. p = 1.8 × 10^{-12}). Notably, for EVP cells, we observed an epithelial-fibroblast phenotype, such as normal human dermal fibroblast cells (Z score = -1.68; adj. p = 1.76 × 10^{-41}) and normal human bronchial epithelial cells (Z score = -1.64; adj. p = 1.34 × 10^{-12}). The expression of low levels of hemoglobin genes in most endothelial cells might be related to the sensitivity of the method and can also be identified in previous
endothelial-related scRNA datasets (Emara et al., 2014; Newton et al., 2006; Saha et al., 2014, 2017; Zhao et al., 2018).

Validation of Subpopulation Clusters One and Two

Flow cytometry analysis was conducted to demonstrate that cluster one (D) and two (EVP) subpopulations fit our proposed endothelial hierarchy based on CD31 and VEGFR2 levels (Figure 4A). To validate the gene expression findings detailed earlier, we fluorescence-activated cell sorting (FACS)-sorted EVP and D cells from the aorta as demonstrated in Figure 4A. Through immunofluorescence, we show that markers, such as decorin (Dcn) and periostin (Postn), had higher expression in EVP cells compared to D at the protein level, as suggested by the scRNA-seq data (Figure 4B).

Figure 3. Gene Co-expression Networks and Correlation Analysis

(A and B) Weighted gene co-expression analysis of D (A) and EVP (B) populations revealed two regulatory modules in each, which are shown as orange and blue. (C–F) The genes from each module were analyzed for significantly enriched biological processes using gene ontology annotations for the D population (C and E) and the EVP population (D and F). The size of the circles in (C)–(F) represents the number of significantly enriched genes per module in each process. The adjusted p value (false discovery rate) threshold was 0.05, and significance is shown as a gradient of color in each circle from blue (less significant) to red (more significant).
A redundant finding in our gene expression analysis of EVP cells was the group of genes involved in metabolism. We therefore explored cell metabolism by comparing mitochondrial activity using flow cytometry and immunofluorescence between the populations EVP and D. Using tetramethylrhodamine methyl ester (TMRM), a marker of mitochondrial membrane potential and a gold standard in measuring mitochondrial activity (Scaduto and Grotyohann, 1999), we clearly demonstrated that the EVP population had significantly higher mitochondrial activity compared to D, based on TMRM staining under immunofluorescence Z stack (Figure 4C) and geometric mean fluorescence intensity (MFI) (44,669 ± 2,354 EVP versus 29,732 ± 543 D; p < 0.05; n = 3; Figures 4D and 4E). On confocal microscopy in sorted cells, we could clearly observe that the increased TMRM staining was related to increased number of mitochondria (Figure 4C). In addition, within our scRNA-seq and bulk RNA-seq datasets, we observed a 2.9- and 3.0-fold increase, respectively, of Cyb5a (mitochondrial membrane protein) expression in EVP cells compared to D, further validating the notion of higher mitochondrial content within the EVP population (Figure 4F). These changes in mitochondrial number and activity are likely to reflect variation in metabolic activity in EVP compared to D cells and validate the scRNA gene expression. Overall, these findings validate key significant changes in gene expression between these two populations of endothelial cells, highlighting different properties in mitochondrial activity, adhesion, and interaction with extracellular matrix proteins.

**Cell Cycle Phase Classification**

EVP cells are more likely to be a quiescent population compared to a more actively proliferating population, such as the D cells (Patel et al., 2017), and the prediction of cell cycle phases can help to stratify the cell types. To classify the cell cycle phase of each individual cell, we used the cyclone method to calculate the G1, G2M, and S scores for each cell using a machine learning dataset trained on the mouse cell cycle (Lun et al., 2016; Scialdone et al., 2015). Our data revealed that the vast majority of cells (95.2%) exhibited a G1 quiescent phenotype, 4% were classified as G2M, and 0.8% were classified as S phase. We asked whether specific clusters were more quiescent or proliferative than others and observed an enrichment of proliferating cells (G2M) in clusters one (D: 5.9%; p = 1.83 × 10^{-16} [Pearson’s chi-square test]) and three (D': 19.5%; p = 2.86 × 10^{-6}) compared to the EVP population (0.7%) in cluster two. Furthermore, only 0.2% of EVP cells were in S phase, indicating these cells are more quiescent than the D (1.1%; p = 6.15 × 10^{-5}) or D' (4.3%; p = 1.52 × 10^{-3}).
Cell Fate Trajectory Analysis of Vascular Progenitors

We previously showed that EVP and D cells in whole tissue exist as a continuum of differentiation states. To investigate the transition of the EVP cells to the terminally differentiated D state at single-cell resolution, we performed a cell fate trajectory and pseudotime analysis using the Monocle 2 software (Qiu et al., 2017) for cells in clusters 1–3. Cluster four, which we classified as muscle cells, was excluded from this analysis, allowing us to focus on endothelial cells only. Our trajectory analysis revealed a continuum of cells with three distinct branch points, showing a root corresponding predominantly to the EVP population (cluster two) and a terminal population corresponding to D cells (cluster one; Figures 5A and 5B). States 1, 2, 3, and 7, comprised mainly of EVP cells, also contained a smaller proportion of D cells (Figures 5B and 5C; Table 2). These less differentiated D cells may represent the “transit-amplifying” (TA) population identified in our previous study (Patel et al., 2017). We observed a branch point arising mainly from cluster one (state 4; Figure 5C) that split into two branches: one comprised of D cells (state 5) and another comprised of cells from clusters one and three (state 6). To determine whether the individual states represent important subpopulations within the D and EVP populations, we tested for key gene signatures using differential expression between pairs of states. However, this did not reveal sufficiently distinct signatures to allow robust cell subtype classification beyond the root (EVP) and tip (D and D’) classifications.

Furthermore, when we overlaid the cell cycle phase information for each individual cell onto the trajectory tree, we observed a higher proportion of cells in G2M in trajectory states corresponding to more differentiated cells. This finding is consistent with our initial cell cycle analysis of each cluster. No batch effects were observed between the three pooled libraries (Figures S6C–S6E).

The high resolution of scRNA-seq enabled us to estimate the proportion of cells with the potential to transition from one subpopulation to another. The transition scores for each, representing the percentage of cells with transcriptional potential to transition, reveal predominantly stable populations at each stage of differentiation.

DISCUSSION

Endothelial heterogeneity is a more and more accepted hallmark of blood vessel biology, with subpopulations identified across or
within vascular beds. Despite major progress in determining the biology of these populations, an unbiased approach to delineating the endothelial heterogeneity has not yet been undertaken. To better understand this heterogeneity during homeostasis in adults, we here studied the endothelium of the largest arterial structure, the aorta, via scRNA-seq. We identified three endothelial populations, one suggestive of a progenitor population that displayed a more mesenchymal phenotype, quiescence, and high mitochondrial content, and which preceded two mature differentiated endothelial populations in pseudotime trajectories. These findings shed light on the homeostasis of the vascular endothelium.

Previous studies have reported on single-cell sequencing efforts, including endothelial cells in cancer (Sun et al., 2017) or from whole organ (e.g., heart) cluster analysis (Skelly et al., 2018). However, to date, a detailed analysis of homeostatic endothelium is lacking. We previously reported an endothelial hierarchy in the adult vasculature that is governed by an EVP that can give rise to a differentiated D population (Patel et al., 2017). This relationship between the endothelial subpopulations has been substantiated by in vivo lineage tracing and functional assays demonstrating that EVP cells have self-renewal and engraftment capacity. In this previous work, EVP and D cells were initially defined through observation of levels of common surface markers, such as CD31 and VEGFR2. The unbiased nature of the present study further strengthens these previous findings. We observed extremely high and significant correlation of gene expression between EVP and D and the respective clusters of endothelial cells identified here, strongly indicating that these are representations of the same populations. Sequencing depth and average read per cell reached saturation, ensuring that the observed heterogeneity reflected biological phenomena (Figure S1). Many of the genes upregulated within cluster two, that we termed the EVP, are identified as being crucial in maintaining stem cell function, phenotype, and quiescence, such as Sox9 and Il33 and the cyclin-dependent kinase inhibitor family (Marqués-Torrejón et al., 2013; Patel et al., 2017; Scott et al., 2010). Furthermore, metabolic activity has been demonstrated to be crucial in maintaining the self-renewal capacity of stem cell populations (Ito and Suda, 2014; Khacho and Slack, 2017; Simsek et al., 2010; Takubo et al., 2013). Within cluster two, metabolic activity was upregulated, and this population displayed a specific increase in expression of genes involved in mitochondrial activity, as validated by measuring TMRM levels as a surrogate marker of mitochondrial activity, further validating a major hierarchical difference between EVP and D cells. Our analysis also revealed in a small proportion of D cells the potential to transition in the reverse direction to EVP. However, there is little in vivo evidence in the context of wound healing supporting the reversion of D cells to EVP. It is more likely that these cells represent the transit-amplifying population we previously identified (Patel et al., 2017) and that the expression levels of the cell surface proteins used to identify them by flow cytometry are not strongly correlated with their mRNA expression. Fate tracing using specific genes from these transitional stages would definitively identify their role in the endothelial hierarchy.

Of note, despite the high level of correlation in gene expression between EVP and cluster two or D and cluster one, many key genes initially identified by us as determinant in these populations could not be found. Indeed, the LASSO machine learning identified a set of markers for these populations based on the single-cell data. This might be related to the preferential sequencing of highly expressed genes in scRNA-seq. It is likely that this is due to a lack of sequencing depth, because the dataset was sequenced to an average depth of 53,631 reads/cell and QC metrics show “sequencing saturation” and “median genes per cell” approached maximum levels (Figure S1). Similarly, other groups have also proposed the existence of resident “vascular stem cell populations.” These have been identified using cell surface markers, such as CD157 in the liver (Wakabayashi et al., 2016) or activated protein C receptor in mammary tissue (Yu et al., 2016). However, to our surprise, neither of these markers could be robustly identified in our single-cell dataset and distinguish populations with the same clarity as Il33, Sox9, or Sox18. Certainly, further studies are warranted to identify the expression of these stem cell markers within aortic endothelial clusters.

Another key finding was the homogeneous high expression of Pdgfra in EVPs. This is a well-described cell surface receptor that accurately distinguishes mesenchymal stem cell populations from both mouse and human tissue of various organs (Houlihan et al., 2012; Uezumi et al., 2014). The identification of Pdgfra within cluster two in addition to a large number of extracellular matrix genes raises the potential for this population having bi-potential capability, a capacity to give rise to both endothelial and mesenchymal cells. We recently reported such bi-potential progenitor within the endothelium of the human term placenta, isolated from the vascular bed (expressing VE-cadherin) but also having high expression of mesenchymal genes, including Pdgfra, and giving rise to both endothelial and mesenchymal colonies in vitro (Shafiee et al., 2018). An additional study during embryonic development demonstrated the crucial role for Pdgfra within the endothelium in forming mesenchyme-derived structures, such as cardiac ventricles (Aghajanian et al., 2017). The resultant loss of Pdgfra in the developing endothelium prevented the correct formation of ventricle structures. Therefore, this further suggests that cluster two/EVP may present as a conserved bi-potential cell in the adult endothelium that maintains this exclusive capacity and warrants further study in vivo.

In summary, we report an unbiased single-cell transcriptomic approach in delineating subpopulations of the homeostatic

| Table 2. Proportion of Cells in Each “State” as Determined by Pseudotime Analysis in Monocle 2 |
|---------------------------------|---|---|---|---|
| State  | Cluster | Cluster | Cluster | Cluster |
| 1      | 6.2  | 17.3 | 0  | 0  |
| 2      | 2.7  | 3.3  | 0  | 0  |
| 3      | 4.0  | 5.6  | 0  | 0  |
| 4      | 6.8  | 2.0  | 0  | 0  |
| 5      | 19.1 | 0  | 0  | 0  |
| 6      | 9.3  | 0.5  | 2.7 | 1.3 |
| 7      | 4.8  | 14.5 | 0  | 0  |
endothelial compartment from the adult aorta. The identification of two main clusters very strongly recapitulates recently identified progenitor EVP and differentiated D populations. This provides an opportunity to understand the molecular events occurring during the transition from progenitor to differentiated cell in the resting endothelium of the adult aorta.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
- METHOD DETAILS
  - Tissue processing of murine aorta and FACS
  - Immunofluorescence and Microscopy
  - Metabolism Assay
  - Single-cell sequencing
- QUANTIFICATION AND STATISTICAL ANALYSIS
  - Linear modeling of transcript expression
  - Cell fate trajectory analysis
  - Gene co-expression analysis
  - Correlation analysis
- DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.celrep.2019.04.102.

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AUTHOR CONTRIBUTIONS

S.W.L. performed the scRNA-seq and computational analysis and wrote and edited the manuscript. J.P. generated cells for scRNA-seq, completed all biological validation and analysis, and wrote and edited the manuscript. S.B.A. assisted in scRNA-seq, S.-L.S. assisted in biological validation and FACS. H.Y.W. assisted in biological validation and microscopy. J.T. assisted in biological validation and metabolism assay. I.W. assisted in data analysis and editing of manuscript. J.E.P. assisted in computational analysis of scRNA-seq data and edited the manuscript. K.K. conceived and supervised the study, wrote and edited the manuscript, and generated funding.

DECLARATION OF INTERESTS

J.P. and K.K. are co-inventors relating to the isolation of endothelial progenitors from the placenta.

REFERENCES

Adams, R.H., and Alitalo, K. (2007). Molecular regulation of angiogenesis and lymphangiogenesis. Nat. Rev. Mol. Cell Biol. 8, 464–478.
Aghajanian, H., Cho, Y.K., Rizer, N.W., Wang, G., Li, L., Degenhardt, K., and Jain, R. (2017). Pdgfrα functions in endothelial-derived cells to regulate neural crest cells and the development of the great arteries. Dis. Model. Mech. 10, 1101–1108.
Aird, W.C. (2007a). Phenotypic heterogeneity of the endothelium: I. Structure, function, and mechanisms. Circ. Res. 100, 158–173.
Aird, W.C. (2007b). Phenotypic heterogeneity of the endothelium: II. Representative vascular beds. Circ. Res. 100, 174–190.
Aird, W.C. (2012). Endothelial cell heterogeneity. Cold Spring Harb. Perspect. Med. 2, a006429.
Anders, S., and Huber, W. (2010). Differential expression analysis for sequence count data. Genome Biol. 11, R106.
Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. J. Roy. Stat. Soc. B Met. 57, 289–300.
Chen, E.Y., Tan, C.M., Kou, Y., Duan, Q., Wang, Z., Meirelles, G.V., Clark, N.R., and Ma’ayan, A. (2013). Enrichr: interactive and collaborative HTML5 gene list enrichment analysis tool. BMC Bioinformatics 14, 128.
Dobin, A., Davis, C.A., Schlesinger, F., Drenkow, J., Zaleski, C., Jha, S., Batut, P., Chaisson, M., and Gingeras, T.R. (2013). STAR: ultrafast universal RNA-seq aligner. Bioinformatics 29, 15–21.
Donovan, P., Patel, J., Dight, J., Wong, H.Y., Sim, S.-L., Murigneux, V., Francois, M., and Khorshetehtani, K. (2019). Endovascular progenitors infiltrate melanomas and differentiate towards a variety of vascular beds promoting tumor metastasis. Nat. Commun. 10, 18.
Eichmann, A., Yuan, L., Moyon, D., Lenoble, F., Pardanaud, L., and Breant, C. (2005). Vascular development: from precursor cells to branched arterial and venous networks. Int. J. Dev. Biol. 49, 259–267.
Emara, M., Turner, A.R., and Allainou-Turner, J. (2014). Hypoxia differentially upregulates the expression of embryonic, fetal and adult hemoglobin in human glioblastoma cells. Int. J. Oncol. 44, 950–958.
Fang, S., Wei, J., Pentimikko, N., Leinonen, H., and Salven, P. (2012). Generation of functional blood vessels from a single c-kit+ adult vascular endothelial stem cell. PLoS Biol. 10, e1001407.
Herbert, S.P., and Stanier, D.Y. (2011). Molecular control of endothelial cell behaviour during blood vessel morphogenesis. Nat. Rev. Mol. Cell Biol. 12, 551–564.
Houlihan, D.D., Mabuchi, Y., Morikawa, S., Niibe, K., Araki, D., Suzuki, S., Okano, H., and Matsuzaki, Y. (2012). Isolation of mouse mesenchymal stem cells on the basis of expression of Sca-1 and PDGFR-α. Nat. Protoc. 7, 2103–2111.
Iruela-Arispe, M.L., and Davis, G.E. (2009). Cellular and molecular mechanisms of vascular lumen formation. Dev. Cell 16, 222–231.
Ito, K., and Suda, T. (2014). Metabolic requirements for the maintenance of self-renewing stem cells. Nat. Rev. Mol. Cell Biol. 15, 243–256.
Khacho, M., and Slack, R.S. (2017). Mitochondrial activity in the regulation of stem cell self-renewal and differentiation. Curr. Opin. Cell Biol. 49, 1–8.
Langfelder, P., and Horvath, S. (2008). WGCNA: an R package for weighted correlation network analysis. BMC Bioinformatics 9, 559.
Lun, A.T., McCarthy, D.J., and Marioni, J.C. (2016). A step-by-step workflow for low-level analysis of single-cell RNA-seq data with Bioconductor. F1000Res. 5, 2122.
Marqués-Torrejón, M.A., Porlan, E., Banito, A., Gómez-Ibáñez, E., Lopez-Contreras, A.J., Fernández-Capetillo, O., Vidal, A., Gil, J., Torres, J., and Farías, I. (2013). Cyclin-dependent kinase inhibitor p21 controls adult neural stem cell expansion by regulating Sox2 gene expression. Cell Stem Cell 12, 88–100.
Newton, D.A., Rao, K.M., Dluhy, R.A., and Baatz, J.E. (2006). Hemoglobin is expressed by alveolar epithelial cells. J. Biol. Chem. 281, 5668–5676.

Nguyen, Q.H., Lukowski, S.W., Chiou, H.S., Senabouth, A., Bruxner, T.J.C., Christ, A.N., Paipint, N.J., and Powell, J.E. (2018). Single-cell RNA-seq of human induced pluripotent stem cells reveals cellular heterogeneity and cell state transitions between subpopulations. Genome Res. 28, 1053–1066.

Patel, J., Seppanen, E.J., Rodero, M.P., Wong, H.Y., Donovan, P., Neufeld, Z., Fisk, N.M., Francois, M., and Khotabeh, K. (2017). Functional definition of progenitors versus mature endothelial cells reveals key Sox5-dependent differentiation process. Circulation 135, 786–805.

Qiu, X., Mao, Q., Tang, Y., Wang, L., Chawla, R., Pliner, H.A., and Trapnell, C. (2017). Reversed graph embedding resolves complex single-cell trajectories. Nat. Methods 14, 979–982.

Shafiee, A., Patel, J., Hutmacher, D.W., Fisk, N.M., and Khosrotehrani, K. (2018). Meso-endothelial bipotent progenitors from human placenta display distinct molecular and cellular identity. Stem Cell Reports 10, 890–904.

Simsek, T., Kocabas, F., Zheng, J., Deberardinis, R.J., Mahmoud, A.I., Olson, E.N., Schneider, J.W., Zhang, C.C., and Sadek, H.A. (2010). The distinct metabolic profile of hematopoietic stem cells reflects their location in a hypoxic niche. Cell Stem Cell 7, 380–390.

Skelly, D.A., Squiers, G.T., McLeellan, M.A., Bolisetty, M.T., Rosson, P., Rosenthal, N.A., and Pinto, A.R. (2018). Single-cell transcriptional profiling reveals cellular diversity and intercommunication in the mouse heart. Cell Rep. 22, 600–610.

Sun, Z., Wang, C.Y., Lawson, D.A., Kwek, S., Velozo, H.G., Owyong, M., Lai, M.D., Cong, L., Wilson, M., Su, H., et al. (2017). Single-cell RNA sequencing reveals gene expression signatures of breast cancer-associated endothelial cells. Oncotarget 9, 10945–10961.

Szklarczyk, D., Franceschini, A., Wyder, S., Forsalund, K., Heller, D., Huerta-Cepas, J., Simonovic, M., Roth, A., Santos, A., Tsafou, K.P., et al. (2015). STRING v10: protein-protein interaction networks, integrated over the tree of life. Nucleic Acids Res. 43, D447–D452.

Takubo, K., Nagamatsu, G., Kobayashi, C.I., Nakamura-Ishizu, A., Kobayashi, H., Ikeda, E., Goda, N., Rahimi, Y., Johnson, R.S., Soga, T., et al. (2013). Regulation of glycosylation by Pdk functions as a metabolic checkpoint for cell cycle quiescence in hematopoietic stem cells. Cell Stem Cell 12, 49–61.

Uezumi, A., Fukada, S., Yamamoto, N., Ikimoto-Uezumi, M., Nakatani, M., Morita, M., Yamaguchi, A., Yamada, H., Nishino, I., Hamada, Y., and Tsuchida, K. (2014). Identification and characterization of PDGFRα+ mesenchymal progenitors in human skeletal muscle. Cell Death Dis. 5, e1186.

Wakabayashi, T., Naito, H., Suenohru, J.I., Lin, Y., Kawai, H., Iba, T., Kouno, T., Ishikawa-Kato, S., Furuno, M., Takara, K., et al. (2018). CD157 marks tissue-resident endothelial stem cells with homeostatic and regenerative properties. Cell Stem Cell 22, 384–397.e6.

Yu, Q.C., Song, W., Wang, D., and Zeng, Y.A. (2016). Identification of blood vascular endothelial stem cells by the expression of protein C receptor. Cell Res. 26, 1079–1098.

Zhao, Q., Eichten, A., Parveen, A., Adler, C., Huang, Y., Wang, W., Ding, Y., Adler, N., Nevin, N., T., N., M., et al. (2018). Single-cell transcriptome analyses reveal endothelial cell heterogeneity in tumors and changes following antiangiogenic treatment. Cancer Res. 78, 2370–2382.
STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| PE Rat Anti-Mouse VEGFR2 | Becton Dickinson | 555308; RRID:AB_395271 |
| PE-Cy7 Rat Anti-Mouse CD31 | Becton Dickinson | 561410; RRID:AB_10612003 |
| Alexa Fluor 647 Rat Anti-Mouse CD34 | Becton Dickinson | 560230; RRID:AB_1645200 |
| BV450 Rat Anti-Mouse Lineage Cocktail | BioLegend | 133310; RRID:AB_10768377 |
| Alexa Fluor 488 Rat Anti-Mouse CD144 | eBioscience | 5016730; RRID:AB_10597442 |
| Goat Polyclonal Anti-Decorin | R&D | AF1060; RRID:AB_2090386 |
| Rabbit Polyclonal Anti-Periostin | Abcam | ab14041; RRID:AB_2299859 |
| Donkey Anti-Goat Alexa Fluor 488 | Invitrogen | ab150129; RRID:AB_2687506 |
| Donkey Anti-Rabbit Alexa Fluor 488 | Invitrogen | ab150073; RRID:AB_2636877 |
| Tetramethylrhodamine, Methyl Ester | Life Technologies | T668 |
| Chemicals, Peptides, and Recombinant Proteins | | |
| Collagenase Type I | GIBCO | 17100017 |
| Dispase II | GIBCO | 17105041 |
| Deoxyribonuclease I | Sigma-Aldrich | D5025-375KU |
| Critical Commercial Assays | | |
| Chromium Single Cell 3’ Gel Bead and Library Kit v2 | 10X Genomics | PN-120237 |
| KAPA Library Quantification Kit | Roche | KR0405 – v8.17 |
| Deposited Data | | |
| scRNA-seq data | This paper | E-MTAB-7149 |
| Bulk RNA-seq data | Donovan et al., 2019 | E-MTAB-7148 |
| Experimental Models: Organisms/Strains | | |
| Mouse: C57BL/6JArc | Animal Resources Centre Perth | JAX: 000664 |
| Software and Algorithms | | |
| Cellranger | 10x Genomics | 2.1.0 |
| Ascend | https://github.com/IMB-Computational-Genomics-Lab/ascend | 0.5 |
| Scran | Lun et al., 2016 | 1.8.4 |
| DESeq | Anders and Huber, 2010 | 1.32.0 |
| Monocle2 | Qi et al., 2017 | 2.8.0 |
| scGPS | https://github.com/IMB-Computational-Genomics-Lab/scGPS | 0.9.9 |
| WGCNA | Langfelder and Horvath, 2008 | 1.63 |
| FV31S-SW | Olympus | Ver2.3.1 |
| ArrayExpress | https://www.ebi.ac.uk/arrayexpress | Accession numbers E-MTAB-7149 and E-MTAB-7148 |

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Professor Kiarash Khosrotehrani (k.khosrotehrani@uq.edu.au).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

All mice were treated in accordance with University of Queensland ethics approvals and guidelines for care of experimental animals. C57BL/6 mice (WT) females were obtained from the Animal Resources Centre (Perth, Western Australia).
**METHOD DETAILS**

**Tissue processing of murine aorta and FACS**

Aortas were dissected and digested for 30 mins at 37°C in 1mg/ml collagenase I (GIBCO, Life Technologies, NY, USA), 1mg/ml Dispase (GIBCO, Life Technologies, NY, USA), 150 μg/ml DNase-I (Sigma-Aldrich, St Louis, MO, USA) before passing the suspension through a 70 μm cell strainer. Dissociated single cells in PBS/BSA/EDTA were then incubated with various antibody combinations for multi-parameter flow acquisition and analysis. The following combinations of antibodies were used to assess the endothelial hierarchy populations: Rat anti-mouse VEGFR2 PE, CD31 PE-Cy7 and CD34 Alexa647 (Becton Dickinson, NJ, USA), Rat anti-mouse Lineage cocktail BV450 (Biolegend), Rat anti-mouse CD144 FITC (eBioscience). Fluorescence-activated cell sorting (FACS) was performed on a FACSARia cell sorter (Becton Dickinson, Franklin Lakes, NJ, USA). Doublets were gated out prior to isolation. Isolated cells were then cytospun onto Superfrost Plus Microscopy slides (Thermo Scientific, MA, USA) for immunofluorescence analysis.

**Immunofluorescence and Microscopy**

For this study, primary antibodies used included goat anti-decorin (1:50) and rabbit anti-periostin (1:100) (all from Abcam, MA, USA). Excess and unbound antibody was then removed with 3x 5mins washes in a solution containing 1x PBS/0.1% Tween-20 (Amresco, Solon, Ohio, USA). Secondary antibody conjugated with Alexa Fluor 488 (Invitrogen, Carlsbad, CA, USA) was used for fluorescence detection. Briefly, slides were incubated with secondary antibodies for 40mins at room temperature. Excess antibody was removed by 3x washes in PBS/0.1% Tween-20. Nuclear staining was revealed in specimens mounted with ProLong® Gold mounting media containing DAPI (Invitrogen, Carlsbad, CA, USA). Confocal images were acquired with a Zeiss LSM 710 microscope. Images were obtained at 63x.

**Metabolism Assay**

Filtered aortic cell suspensions were incubated with 20 nM tetramethylrhodamine, methyl ester (TMRM, Thermofisher) for 20 min at 37°C in the dark with agitation. Cells were then washed in ice-cold PBS/BSA/EDTA. Cells were then stained with antibodies against cell surface markers for 30 min in the dark on ice. Excess antibodies were washed off with PBS/BSA/EDTA buffer. The fluorescence of mitochondrial bound TMRM was determined by flow cytometry and immunofluorescence, with excitation at 561 nm and emission at 585 ± 21 nm. Cell populations were analyzed using CytoFLEX S analyzer (Beckman Coulter) followed by further analyses using FlowJo v7-10 software (TreeStar).

**Single-cell sequencing**

FACS-sorted single-cell suspensions were used to generate barcoded single-cell 3’ cDNA libraries for each sample pool with the Chromium Single Cell 3’ Gel Bead and Library Kit v2 (10x Genomics). Libraries were assessed with an Agilent BioAnalyzer High Sensitivity DNA chip, pooled, and quantified with qPCR (KAPA Library Quantification Kit). Denatured libraries were loaded onto an Illumina NextSeq-500 and sequenced using a 150-cycle High-Output Kit to an average depth of 53,631 reads/cell. To process the sequencing data, we used the 10x Genomics cellranger pipeline (v2.1.0), comprising the mkfastq, count and aggr stages. Using cellranger mkfastq, raw base call files were demultiplexed into sample-specific FASTQ files. Sample-specific FASTQ files were quality controlled and aligned to the mouse reference genome (mm10) using STAR aligner (Dobin et al., 2013) during the cellranger count stage. Aligned reads were filtered for valid cell barcodes and unique molecular identifiers. An aggregated, between-sample normalized gene expression matrix was generated using cellranger aggr.

The aggregated single-cell gene expression data generated by cellranger was used as the input for the ascend analysis pipeline (v0.5; https://github.com/IMB-Computational-Genomics-Lab/ascend). Expression levels for each transcript were determined using the number of unique molecular identifiers (UMI) assigned to the transcript. Quality control and filtering steps were performed to remove outlier cells and genes. Cells were excluded if the library size, or number of expressed genes exceeded 3 median absolute deviations, or if mitochondrial or ribosomal reads accounted for more than 20% or 50% of the reads respectively. The relative log expression (RLE) method was used to normalize cell-cell expression variation, and ribosomal and mitochondrial control genes were subsequently excluded from the analysis.

Principal component analysis was performed on the filtered and normalized gene expression matrix, and the first 20 PCs that explained the majority of variance in the data were retained. Unsupervised hierarchical clustering of individual cells was applied using the method previously described (Nguyen et al., 2018). To visualize the patterns of gene expression in each cell in each cluster, t-distributed stochastic neighbor (t-SNE) projections were generated using the PCA-reduced data.

Differential expression analysis was performed with DESeq (Anders and Huber, 2010) in ascend between (i) each cluster compared to the remaining cells, or (ii) between pairs of specific clusters. Differences in gene expression were considered significant if the adjusted P value was below the multiple-testing threshold of 0.01 (Benjamini and Hochberg, 1995) and the absolute log2 expression fold change was ≥ 2.

Cell cycle phase classification of each cell (n = 6,111) was performed using the cyclone function implemented in the scran package (Lun et al., 2016), with a dataset trained on mouse cell cycle gene expression.
QUANTIFICATION AND STATISTICAL ANALYSIS

Unless otherwise noted, all data are represented as mean ± standard error of mean (SEM). Indicated sample sizes (n) represent biological replicates including individual samples, these have been listed in the results section and figure legends where appropriate. Statistical significance where appropriate was determined in GraphPad Prism 7 software by using the Mann-Whitney t-test (non-parametric). For validation of sequencing data, a significance threshold of p < 0.05 was used.

Linear modeling of transcript expression
Linear regression was used to test the expression of each individual gene (n = 17,323) against the cluster classification of the 5,867 cells comprising clusters 1 and 2 only. The overall model P value was adjusted for multiple testing using a Bonferroni threshold calculated as 0.05/17,323 = 2.89x10^{-6} and transcripts with an adjusted P value below the threshold were considered statistically significant.

Cell fate trajectory analysis
Monocle2 (Qiu et al., 2017) was used to order cells in pseudo-time based on the top 500 genes with the highest variation in expression across the 6,031 cells comprising clusters 1-3. The most variable genes were determined using a \log_2(counts + 1)-transformed dataset. The LASSO regression procedure in scGPS (https://github.com/IMB-Computational-Genomics-Lab/scGPS) was used to calculate the deviance coefficient for each of the genes defining each cluster. For the scGPS input, we used the significant differentially expressed genes that were obtained by comparing cluster x to the remaining clusters, a vector of cluster information for each cell, and \log_2(counts + 1)-transformed expression data.

Gene co-expression analysis
To establish gene co-expression networks underlying the EVP and D populations in our single-cell data, we applied weighted gene co-expression network analysis as implemented in the WGCNA package for R (Langfelder and Horvath, 2008).

Correlation analysis
Mean expression values for scRNA-seq clusters and bulk RNA-seq samples was used to perform pairwise Spearman’s correlations. Correlations were performed with known gene markers for EVP and D cells derived from previous studies on normal (Patel et al., 2017) and tumor-derived populations (Donovan et al., 2019), and these were merged with differentially expressed genes defining the scRNA-seq clusters. This yielded a common set of 2,105 genes between all samples that were taken forward for pairwise correlation analysis.

DATA AND SOFTWARE AVAILABILITY

Single-cell RNA sequencing data was deposited in ArrayExpress (https://www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-7149 and bulk RNA sequencing data under accession number E-MTAB-7148.
Supplemental Information

Single-Cell Transcriptional Profiling of Aortic Endothelium Identifies a Hierarchy from Endovascular Progenitors to Differentiated Cells

Samuel W. Lukowski, Jatin Patel, Stacey B. Andersen, Seen-Ling Sim, Ho Yi Wong, Joshua Tay, Ingrid Winkler, Joseph E. Powell, and Kiarash Khosrotehrani
Transcriptional profiling of aortic endovascular progenitors at single cell resolution

Lukowski and Patel et al – Cell Reports
Supplementary Results

Sample 1

Figure S1: Cellranger sequencing metrics for samples 1-3. Sequencing Saturation (left) and Median Genes Per Cell (right) compared to Mean Read Depth Per Cell. Related to Figure 1.
Figure S2: STRING network analysis of genes in the D blue module detected by WGCNA. Related to Figure 3.

Enriched pathways: Immune system; positive regulation of metabolic process; cell adhesion
**Figure S3**: STRING network analysis of genes in the D orange module detected by WGCNA. Related to Figure 3.
**Enriched pathways:**
Metabolic process; oxidative phosphorylation

**Figure S4:** STRING network analysis of genes in the EVP blue module detected by WGCNA. Related to Figure 3.
**Figure S5:** STRING network analysis of genes in the EVP orange module detected by WGCNA. Related to Figure 3.

**Enriched pathways:**
Metabolic process; vasculature development
**Figure S6:** Cell cycle classifications shown as stacked barplots for each batch (A) and each cluster (B). (C) Cell cycle analysis scores for each individual cell were projected onto the monocle trajectory analysis. tSNE plot showing overlay of the three sample libraries coloured by batch. No obvious batch effects were observed (D). Panel (E) shows cells on the trajectory plot separated by cell cycle stage. Related to Figure 5.