Gene Expression Within a Human Choroidal Neovascular Membrane Using Spatial Transcriptomics

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METHODS. To study how gene expression is altered in focal areas of pathology, we performed spatial RNA sequencing on a human donor eye with macular neovascularization as well as a healthy control donor. We performed differential expression to identify genes enriched within the area of macular neovascularization and used deconvolution algorithms to predict the originating cell type of these dysregulated genes.

RESULTS. Within the area of neovascularization, endothelial cells demonstrated increased expression of genes related to Rho family GTPase signaling and integrin signaling. Likewise, VEGF and TGFB1 were identified as potential upstream regulators that could drive the observed gene expression changes produced by endothelial and retinal pigment epithelium cells in the macular neovascularization donor. These spatial gene expression profiles were compared to previous single-cell gene expression experiments in human age-related macular degeneration as well as a model of laser-induced neovascularization in mice. As a secondary aim, we investigated regional gene expression patterns within the macular neural retina and between the macular and peripheral choroid.

CONCLUSIONS. Overall, this study spatially analyzes gene expression across the retina, retinal pigment epithelium, and choroid in health and describes a set of candidate molecules that become dysregulated in macular neovascularization.

Keywords: neovascularization, age-related macular degeneration (AMD), gene expression, choroid, spatial RNA sequencing

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ge-related macular degeneration (AMD) is a common A cause of blindness in which the central retina (termed the macula) degenerates, leading to a loss of high acuity center vision. Advanced stages of AMD include a progressive deterioration (geographic atrophy) of the retinal pigment epithelium and choriocapillaris vascular layer and/or abnormal new blood vessel growth called macular neovascularization (MNV). In MNV, pathologic angiogenesis can result in severe and often rapid vision loss due to retinal damage from hemorrhage and vascular leakage. Thanks in large part to developments in imaging, MNV can be reproducibly categorized as type I (formerly "occult" choroidal neovascularization) that arises from the choroidal vasculature and lies between the retinal pigment epithelium (RPE) and Bruch's membrane, type II (formerly "classic" choroidal neovascularization) that similarly arises from the choroid but that grows into the subretinal space, and type III MNV that originates

from the retinal vasculature rather than the choroid.¹ MNV can be exudative, with serous fluid seeping past the compromised RPE and damaging the overlying photoreceptors. By contrast, non-exudative MNV has been conjectured to be protective against geographic atrophy by re-establishing a choriocapillaris-like layer to meet the metabolic demands of the retina.²

The application of anti-VEGF agents has been remarkably effective at treating MNV, and millions of patients have had tremendous positive visual outcomes with this therapy.³ In most patients, anti-VEGF is effective at resolving subretinal and intraretinal fluid.⁴ However, in some patients, the response to anti-VEGF drugs may be limited or wear off over time,^{5–8} and the exact mechanism(s) of this attenuated response remain poorly understood.⁹ The visual consequences of MNV and the frequency of associated treatment (often monthly clinic-based injections) signif-

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icantly impacts affected individuals' quality of life.¹⁰ An improved understanding of the pathophysiologic mechanisms involved in MNV may inform the development of targeted interventions for those with limited response to anti-VEGF or those that develop geographic atrophy despite treatment.¹¹

Human neovascular membranes removed during surgery¹² and endothelial cells from the mouse model of laser induced choroidal neovascularization¹³ have been previously studied at the transcriptional level using bulk and single-cell RNA sequencing, respectively. We recently evaluated choroidal gene expression at the single-cell level in a series of eyes from donors with atrophic AMD, age-matched controls, and two eyes with MNV.14 Such approaches highlight dysregulated gene expression in MNV. However, a limitation of transcriptomic studies to date is that the spatial location of each cell is not captured, even when performed at the single cell level. For example, when tissue from an MNV is dissociated, it is not possible to discriminate between endothelial cells within the neovascular membrane versus those in the adjacent and underlying choroid. To detect transcriptomic features within a MNV with greater specificity, spatial analysis of gene expression patterns offers greater potential.

In the current study, we performed spatial RNA sequencing on two human donor eyes, one from an unaffected control and one eye from a patient with type I MNV. Consistent with our expectations, gene expression patterns in the normal retinal outer nuclear, inner nuclear, and ganglion cell layers were highly enriched for photoreceptor, Müller cell, and ganglion cell transcriptomic profiles, respectively. We further identified gene signatures spatially enriched in the neovascular membrane compared to normal choroid, and endothelial cells in this region had increased expression of genes related to Rho family GTPase and integrin signaling. Such spatial profiles were compared to previous single-cell gene expression patterns in human AMD endothelial cells¹⁴ and models of laser-induced neovascular endothelial cells in mice.¹³ As a secondary aim, we identified gene expression patterns that varied spatially within the macular neural retina and compared these results to previous regional single-cell RNA sequencing studies.¹⁵

MATERIALS AND METHODS

Human Donor Eyes

All donor tissue used in this study was recovered by the Iowa Lions Eye Bank with full consent from the next of kin for use in research and evaluation of donor clinical records. All studies were performed in compliance with the Declaration of Helsinki. For the spatial transcriptomics experiments, eyes from two donors were used. The first was from a 78-yearold Caucasian female with a history of cataract removal in both eyes but with an otherwise unremarkable ophthalmic history. The posterior pole of this control donor appeared grossly normal (Fig. 1A). The second was from an 85-yearold Caucasian female with neovascular AMD in both eyes. Visual acuity was 20/20 in the right eye and 20/250 in the left eye, with records of an involuted choroidal neovascular membrane in both eyes for the past 5 years. She had received anti-VEGF agents at some at least 5 years before her passing, and had been monitored for approximately 5 years without change. Approximately a month before her passing from a ruptured aortic aneurysm, she had received



FIGURE 1. Gross, clinical, and histological characterization of eyes used for spatial transcriptomic experiments. Gross photos from the control (A) and MNV (B) donor after dissection. (A) The 4-mm punch was acquired centered on the fovea and approximately 12 mm superotemporal to the macula. Whereas the macula of the healthy control donor appeared grossly normal (A), the MNV donor macula (B) demonstrated a disc-diameter area of elevated vellow material and atrophy. Near infrared scan (C) and optimal coherence tomography scan (D) of the right eye from the patient with macular neovascularization showing a broad, low-lying fibrovascular pigment epithelial detachment with "double-layer sign" (arrowhead). Reticular pseudodrusen (dark rounded profiles on near infrared scan) were noted superior to the lesion along the arcade vessels. Histological macular tissue sections from the control (E) and MNV (F) donor stained with hematoxylin and eosin. Note the thinning of the outer nuclear layer and fibrovascular membrane in the MNV eye. ONL, outer nuclear layer; IS, inner segments; OS, outer segments; NVM, neovascular membrane; BrM, Bruch's membrane; CHO, choroid.

a single injection of anti-VEGF in the OD. From the right eyes of both donors, a 4 mm diameter macular punch was collected centered on the fovea centralis, as was a 4 mm peripheral punch approximately 12 mm superotemporal from the macula. Both central and peripheral punches (spanning the retina, choroid, and sclera) were placed in optimal cutting temperature compound (OCT; Sakura) and were rapidly snap frozen in a liquid nitrogen bath before storage at -80° C.

Spatial RNA Library Generation

Cryostat sections were collected at a thickness of $10 \,\mu\text{m}$ onto a precooled Visium slide (10X Genomics, Pleasanton, CA, USA). Visium slides with the sections were fixed, stained, and imaged with hematoxylin and eosin using a 20X objective on an Olympus BX61 Upright Microscope. Tissue was then permeabilized for 12 minutes, which was established as an optimal permeabilization time based on tissue optimization time-course experiments. The poly-A mRNAs from the sections were released and captured by the poly(dT) primers precoated on the slide, including spatial barcodes and unique molecular identifiers (UMIs). After reverse transcription and second strand synthesis, the amplificated cDNA samples from the Visium slides were transferred, purified, and quantified for library preparation.

Reverse transcription, second strand synthesis and denaturation, and cDNA amplification were performed as described by the manufacturer (10X Genomics Spatial Gene Expression Protocol CG000239 Revision F). The final libraries were pooled and sequenced on the NovaSeq 6000 platform (Illumina, San Diego, CA, USA) generating 100-bp, paired end reads.

Bioinformatic Analysis

FASTQ files were generated from base calls with the bcl2fastq software (Illumina, San Diego CA, USA). FASTQ files were mapped to the human GRCh38 reference genome with Spaceranger (version 1.3.1). Raw FASTQ files and processed data matrices have been deposited in the Gene Expression Omnibus database (GSE234047). Count data from each slide was loaded into Seurat¹⁶ with the Read10X_Image() function. All sections underwent normalization with the SCTransform() function. Data from all sections were merged and subsequently integrated into one dataset before PCA, clustering, and uniform manifold approximation and projection (UMAP) dimensionality reduction.

In addition, we performed a deconvolution analysis with CSIDE²¹ to estimate the proportion of different cell types contributing RNA to each spot using our previous singlecell expression studies in the human retina¹⁵ and RPEchoroid⁴⁹ as references for cell type label transfer. Using the deconvoluted weights, differential expression analysis was completed with the run.CSIDE() function in the spacexr R package (version 2.0.1). To increase the number of candidate genes available for comparison with other datasets, CSIDE was run in the full (non-doublet) mode, the false discovery rate (FDR) was set to 0.9, and the cell type threshold was set to 10. Three sets of differential analyses were completed. First, gene expression was compared between spots overlapping the MNV to spots outside of the MNV or in control tissues. Covariates for the analysis included cell type deconvolution and the percentage of spot area overlapping with the MNV (see below). MNV-enriched genes were compared to previous single-cell RNA sequencing studies of laserinduced choroidal neovascularization in mice (GSE77199¹³) and two human donors with MNV (GSE18332014). Likewise, endothelial and RPE genes with a log_2FC enrichment > 0.5 in the MNV sample were used as input for ingenuity pathway analysis. Second, gene expression was spatially evaluated along a gradient of distance to the fovea. We measured the distance between each spot and the center of the section and scaled this distance from 0 to 1. This distance and cell type deconvolution were used as the covariates for differential expression analysis, and results were compared to a single-cell RNA sequencing study of foveal (1 mm) versus parafoveal (4 mm) human retina (GSE169046¹⁵). Third, we compared gene expression between macular and peripheral choroidal deconvoluted spots. Spots were manually categorized as macular or peripheral prior to differential expression.

Manual Classification of Spots

The LoupeBrowser (10X Genomics, Pleasanton, CA, USA) was used to manually annotate spots originating in macular versus peripheral sections and overlying retinal versus RPE/choroid tissue. These labels were added to the meta data of each section within Seurat. In addition, the MNV was manually traced and intersected with overlying spots, and ImageJ was used to quantify the proportion of each spot that overlapped the neovascular membrane. Spots overlying the sclera were excluded from all analyses.

Immunofluorescence

In order to assess the protein levels and distributions of two genes with altered expression in neovascularization of mouse single cell,¹³ human single cell,¹⁴ and human spatial data (current study), immunohistochemical detection was performed. Paraformaldehyde fixed cryostat sections of the maculae of 3 donors with neovascular membranes (specific clinical subtype not always available) were collected¹⁷ and dual labeled with antibodies directed against TIMP3 and NNMT. Sections were blocked in 1 mg/mL bovine serum albumin (Sigma) in phosphate buffered saline for 15 minutes and incubated with primary antibodies for 1 hour. Antibodies used were rabbit anti-TIMP3 (Thermo Fisher PA5-32621, 1:70 dilution; secondary Invitrogen A21206 AlexaFluor-488 donkey anti-rabbit antibody at 10 ng/µL) and mouse anti-NNMT (Abcam Ab119758, at 10 ng/µL; secondary Invitrogen A10036 AlexaFluor-546 donkey anti-mouse antibody at 10 ng/µL). Sections were rinsed and incubated in the dark with secondary antibody solution for 30 minutes prior to washing and coverslipping. Photomicrographs were collected using identical conditions for regions inside and outside of the neovascular membrane.

RESULTS

Clinical and Histologic Evaluation of Eyes Used in Transcriptomic Experiments

Ocular tissue from two human donors was acquired for spatial transcriptomic experiments. The first donor was a 78-year-old Caucasian woman with a history of cataract removal but no other ocular history. The posterior pole of this control donor appeared grossly normal with normal foveal and choroidal pigmentation (see Fig. 1A). The second donor was an 85-year-old Caucasian woman with a > 5-year year history of MNV in both eyes. A yellow elevation of the retina with areas of both hypopigmentation and hyperpigmentation was observed by gross examination in both maculae of the MNV donor (Fig. 1B, right eye shown). Infrared reflectance imaging and OCT acquired 1 month prior to her death revealed the presence of superior reticular pseudodrusen and a very broad, low-lying fibrovascular pigment epithelial detachment with "double-layer sign"¹⁸ within the macula, consistent with type 1 MNV (Figs. 1C, 1D). Hematoxylin and eosin staining of the control (Fig. 1E) macula demonstrated normal photoreceptor outer segments, RPE, and choriocapillaris. In contrast, the MNV donor had a prominent type I neovascular membrane between Bruch's membrane and the retinal pigment epithelium (Fig. 1F).



FIGURE 2. Spatial RNA sequencing of a control and MNV human donor macula. (**A**) Experimental overview: tissue sections from a control donor (n = 2 sections) and an MNV donor (n = 3 sections) were prepared for spatial RNA sequencing. After fixation and imaging, sections were permeabilized and cDNA libraries were prepared. Hematoxylin and eosin staining of one section of the control donor (**B**). The RNA contribution from retinal, RPE, and choroidal cell types was estimated in each spatially barcoded spot (**C**) and displayed as a pie-chart. Panels (**D**) and (**E**) show a magnified view of the spot area in the macular (*top*) and peripheral (*bottom*) sections. Spots overlying the sclera were excluded from this visualization. (**F**) Uniform manifold approximation and projection (UMAP) dimensionality reduction was applied to visualize clusters of spots with similar gene expression profiles. Each point represents the multidimensional transcriptomic profile from one spot, and points are colored according to the maximally predicted RNA contribution in each spatially barcoded spot. Clustering was visualized according to region (**G**), tissue (**H**), and percentage of the spot area that overlaps with the MNV membrane (**I**).

Spatial RNA Sequencing Overview

A total of five tissue sections (n = 3 MNV and n = 2 control) were acquired for spatial RNA sequencing (Fig. 2A). Macular and peripheral tissues were embedded in the same OCT block and sectioned together onto each capture area, and each section consisted of macular retina, macular RPE/choroid, and peripheral RPE/choroid. The tissue sections were fixed in methanol, stained with hematoxylin and eosin, and imaged (Fig. 2B, Supplementary Fig. S1). Each slide was subsequently permeabilized and cDNA libraries were prepared and sequenced. After mapping to the human genome, unique barcodes were used to spatially link each read to the x/y coordinate of a spot on the gene expression slide.

Consistent with the cell types known to reside in each retinal layer, the different retinal layers demonstrated distinct gene expression patterns. For example, spots in the ganglion cell layer contained high amounts of NEFL, spots overlying the outer nuclear layer were enriched in the conespecific phosphodiesterase PDE6H, and spots overlying the RPE were enriched in LRAT (Supplementary Fig. S2). The diameter of each barcoded spot was 55 µm, which typically covered multiple different cells within the densely cellular retina, RPE, and choroid. To overcome this technical limitation, we next estimated the RNA contribution of different cell types within each spot using a deconvolution algorithm and previously published retinal¹⁵ and choroidal¹⁴ single-cell RNA sequencing studies as reference data sets (Fig. 2C). Indeed, predicted cellular compositions were unique between the different retinal layers (Figs. 2D, 2E), and distinct tiers of spots highly enriched for retinal ganglion cells, bipolar cells, photoreceptors, and RPE cells were observed. Spots with similar multidimensional transcriptomic profiles were clustered and classified based on which cell type was predicted to contribute the most RNA to each spot (Fig. 2F). Clustering was visualized according to region, tissue, and percentage of the spot area that overlapped with the MNV membrane (Figs. 2G, 2H, 2I). As the spots have a relatively large diameter, it is unsurprising that spots often clustered by region (macula versus peripheral; see Fig. 2G) and/or tissue (retina, RPE, and choroid; see Fig. 2H).

Gene Expression Changes in the MNV Lesion

As the primary aim of the study, we investigated gene expression differences between spots within the MNV lesion versus non-MNV RPE and choroid. First, we manually traced the MNV membrane to visualize how the spots overlapped with the pathology (Figs. 3A, 3B, 3C). A total of 144 barcoded spots partially overlapped with the lesion across the 3 sections. However, due to the relatively large spot diameter of 55 microns, no spot was entirely contained in the MNV lesion. We used ImageI to calculate the percentage of spot area overlapping the traced membrane (Fig. 3D), which ranged from < 1% to 91%. Next, we assessed the deconvoluted cell type contributions of spots overlapping the membrane (Fig. 3E). RPE cells were predicted to contribute the most RNA to spots overlapping the MNV, followed by retinal ganglion cells, cone photoreceptor cells, endothelial cells, and pericyte/smooth muscle cells. The presence of inner retinal transcripts within the MNV-overlapping spots most likely represents imprecise deconvolution predictions, as relatively little diffusion of cell-specific transcripts was observed outside of the appropriate cell layers (see Supplementary Fig. S2). However, it is possible that the presence of inner retinal transcripts in the MNV overlapping spots may also be attributed to nonspecific diffusion¹⁹ (which can be due to over permeabilization²⁰) of RNA beyond the cell margins.

Each spot overlying the MNV lesion was composed of multiple underlying cell types and had varying overlap with the neovascular membrane. To account for the multiple sources of variation, we used the CSIDE parametric differential expression algorithm²¹ to identify MNV-enriched genes. CSIDE uses cell type mixtures (identified by deconvolution) as covariates to detect genes whose expression is related to the continuous explanatory variable of spot-MNV overlap area (Fig. 3F). CSIDE then provides differential expression results split for each individual cell type (Supplementary File S1).

Gene expression changes in neovascular endothelial cells and the overlying RPE were of particular interest, and we set out to understand the functional significance of MNV gene expression dysregulation. As such, we performed pathway enrichment analysis to find canonical pathway terms (Figs. 3G, 3H) of genes enriched in MNV-overlapping spots predicted to originate from endothelial cells and RPE cells (see Supplementary File S1). Endothelial genes from spots overlapping the MNV lesion were enriched in pathways terms related to Rho Family GTPase signaling and Integrin signaling (see Fig. 3G). Rho GTPases can regulate angiogenesis by modulating VEGF expression levels.²² The genes CFL1 and ITGB1, which encode the actin binding protein cofilin and the beta-1 integrin subunit, were increased in neovascular endothelial cells. These gene products are involved in Rho signal transduction and have been implicated in endothelial invasion and angiogenesis.^{23,24} Similarly, integrins can stimulate pathological angiogenesis by transducing signaling through VEGF and other growth factors.²⁵ The RAS family genes RHOC (which encodes a protein which colocalizes with integrin $\alpha 5\beta 1$) and RAP1B (which encodes a product that colocalizes with integrin $\alpha v \beta_3$) have been shown to regulate VEGF activation,^{22,26} and both genes were increased in MNV endothelial cells. RPE genes from spots overlapping the membrane were enriched in pathway terms related to ERK/MAPK signaling (see Fig. 3H), which promotes endothelial cell sprouting and angiogenesis.²⁷ For example, MNV-overlapping spots demonstrated increased expression of ITGAV and ITGB8 (two genes highly expressed by the RPE), which encode integrin components that regulate ERK/MAPK signaling and have been associated with angiogenesis and blood vessel network formation.28,29

Similarly, we searched for predicted upstream regulatory factors that could drive the observed gene expression differences in MNV-overlapping spots (Figs. 3I, 3J). Ingenuity pathway analysis identified VEGF and TGFB1 pathways as upstream regulators of MNV gene expression changes predicted to originate from endothelial cells and RPE cells. VEGFs are the central regulators of angiogenesis and are strongly implicated in MNV,³⁰ and the gold standard for treating neovascular AMD is anti-VEGF drugs (e.g. Ref. 31). Likewise, TGFB signaling has complex roles in modulating angiogenesis and vessel formation³² and has been previously shown to participate in choroidal neovascularization in mice.³³ Overall, these data suggest that the observed MNVassociated endothelial and RPE cell gene expression patterns may be driven by known growth factors central to MNV.

Comparing Neovascular Membrane Enriched Genes With Single-Cell Studies

To date, there have been two studies that explored gene expression changes in neovascular choroidal endothelial cells at the single-cell level. First, Rohlenova et al. used a laser injury model to induce choroidal neovascularization in mice before single-cell RNA sequencing of enriched endothelial cells, using the contralateral eye of each mouse as a control.¹³ Second, our group performed single-cell RNA sequencing on 21 human donor choroids, which included 10 control donors and 2 donors with MNV.¹⁴ In both studies, even the pathologic samples contained non-neovascular endothelial cells alongside neovascular endothelial cells. However, the lack of spatial barcoding in these studies makes it difficult to positively identify the endothelial cells within the neovascular membrane and identify unique gene expression features specific to these cells. Despite these



FIGURE 3. Gene expression in a human macular neovascular membrane. (**A**, **B**, **C**) The three macular sections from the MNV donor. The neovascular membrane (*red outline*) is traced in each tissue section and overlapping spot areas (*black circles*) are visualized. Note that the neovascular membrane spanned the entire section in each case. (**D**) The percent area that each spot overlapped the MNV membrane was calculated. A total of 144 spots overlapped the lesion, ranging from <1% to 91% overlap. (**E**) For all spots overlapping the MNV, the relative cell type composition was calculated. (**F**) Design of differential expression analysis. Cell type deconvolution and percent area overlap with the MNV were used as covariates for differential expression analysis. (**G**, **H**, **I**, **J**) After identifying genes enriched in MNV spots, ingenuity pathway analysis was used to identify canonical pathways (**G**, **H**) and upstream regulators (**I**, **J**) of genes predicted to originate from endothelial cells and RPE cells.



FIGURE 4. (**A**) Comparison of MNV-enriched genes in endothelial cells with previous single-cell RNA sequencing studies. Genes upregulated and downregulated in MNV-associated spots according to spatial RNA sequencing (*green*, current study) were compared to a previous single-cell RNA sequencing study of two human donors with MNV (*blue*) and a mouse model of choroidal neovascularization (*red*). Positive \log_2 fold-changes are associated with increased expression in the neovascular endothelial cells. (**B**) Differential expression between macular and peripheral enriched genes in choroidal endothelial cells. Cell type deconvolution and region were used as covariates for differential expression analysis. For endothelial cells, the top differential expression results were compared between this previous study (*purple*) and the current spatial study (*gold*). Positive \log_2 fold-changes are associated with increased expression in the macula.

limitations, we hypothesized that many of the endothelial MNV-enriched genes identified by the current spatial study would also be enriched in the neovascular samples from the previous single-cell investigations despite the known mixed population of normal and pathological endothelial cells.

To test this hypothesis, we identified the top 10 MNV and control enriched genes predicted to originate from endothelial cells in the current study and compared these results to reprocessed single-cell investigations (Fig. 4A). Overall, 70% of single-cell RNA sequencing \log_2 fold-changes were in the same direction as the current spatial study. In the current study, the endothelial gene with the highest MNV \log_2 fold-change was *IL6*. Interleukin 6 is a potent stimulator of vessel sprouting and angiogenesis,³⁴ and *IL6* RNA levels have been previously shown to increase in choroidal macrophages in a laser model of choroidal neovascularization in mice.³⁵ Previous single-cell RNA sequencing data¹⁴ suggests *IL6* is expressed at the RNA level by choroidal endothelial cells, inflammatory macrophages, fibroblasts, and pericytes. Unfortunately, due to the large spot diameter in the current study, it is challenging to distinguish if endothelial cells, the closely associated inflammatory macrophages, or both cell types are responsible for the increased *IL6* within the MNV of this donor.

Regional Gene Expression in the Neural Retina and RPE/Choroid

We next used spatial RNA sequencing to investigate regional patterns of gene expression within the neural retina, RPE, and choroid. We first investigated how gene expression differed between the macular and peripheral choroid of the control donor. Like the MNV differential expression, CSIDE used cell type mixtures (identified by deconvolution) as covariates to detect genes whose expression is related to the binary explanatory variable of macular or peripheral origin (Supplementary File S2). We then compared these spatial enrichments of RPE and endothelial cells to a previous single-cell RNA sequencing study of macular versus peripheral human RPE/choroid.³⁶ We hypothesized that genes previously identified to vary between these two regions in the RPE/choroid would demonstrate similar enrichments in the current spatial dataset (see Fig. 4, Supplementary Fig. S3). A total of 75% of endothelial cell (Fig. 4B) and 75% of RPE cell (Supplementary Fig. S3) log₂ fold-changes were in the same direction as this previous single-cell RNA sequencing study.

Our spatial sections captured macular (but not peripheral) neural retina, so we next investigated how gene expression varies within the macula. Each 4 mm punch used for this investigation was centered on the fovea. Although it is unlikely that the fovea was perfectly centered, we assumed that the center of the section would be closer to the fovea and that the edges of the section would be further from the fovea. Therefore, we measured the distance of each spot to the center of each section, and we scaled this distance from 0 (close to the fovea) to 1 (far from the fovea). Next, we performed differential expression with CSIDE to identify genes with varied regional expression patterns. Similar to the regional choroid investigation, we performed differential expression with CSIDE using cell type mixtures (identified by deconvolution) as covariates to detect genes whose expression is related to the continuous explanatory vari-



FIGURE 5. Comparison of regional retinal gene expression differences between spatial and single-cell RNA sequencing studies. Differential expression was completed to compare genes enriched toward the center and towards the periphery of each section. For macular sections, the distance between each spot and the center of the slide was calculated and scaled from 0 to 1. Cell type deconvolution and distance from the center were used as covariates for differential expression analysis. For cone photoreceptor cells (**A**) and Muller cells (**B**), the top differentially expressed genes between the fovea and parafovea were identified from a previous single-cell RNA sequencing study. Differential expression results were compared between this previous study (*red*) and the current spatial study (*blue*). Positive log₂ fold-changes are associated with increased expression toward the center of the retina.



FIGURE 6. Immunohistochemical detection of TIMP3 (\mathbf{A} , \mathbf{D}) and NNMT (\mathbf{B} , \mathbf{E}) in an eye with a multilayered MNV membrane with subretinal fibrosis separated by a confluent layer of basal laminar deposit (BlamD). Abundant Bruch's membrane (BrM) TIMP3 is present outside of the area of MNV (\mathbf{A}), whereas within the neovascular lesion on the same section there is strong labeling in the BlamD but the signal is attenuated in BrM (\mathbf{D}). Labeling with anti-NNMT antibodies (\mathbf{B} , \mathbf{E}) shows little reactivity outside of the neovascular lesion (\mathbf{B}) but intense binding to elongated cells throughout the MNV membrane (\mathbf{E}). Panels (\mathbf{C}) and (\mathbf{F}) show three channel merged images. GCL, ganglion cell layer; INL, inner nuclear layer; ONL, outer nuclear layer; CHO, choroid. Scalebar = 100 µm.

able of distance from the section center point (Supplementary File S3). We then compared these differential expression results of cone photoreceptor and Müller cells to a previous single-cell investigation of foveal (1 mm) versus parafoveal (4 mm) neural retina from four human donors.¹⁵ We hypothesized that genes previously identified to vary regionally in the neural retina would demonstrate similar enrichments in the current spatial data set (Fig. 5). Overall, 83% of cone photoreceptor and 93% of Müller cell \log_2 fold-changes were in the same direction as the previous single-cell RNA sequencing study. Collectively, these data suggest that even with large spot diameters, spatial RNA sequencing can detect regional-specific gene expression patterns within the heterogenous neural retina and RPE/choroid.

Immunohistochemical Assessment of Two Markers With Differential Expression

The products of two genes that showed differential expression in the mouse CNV model, human sc-RNA-Seq, and the current spatial study (see Fig. 4A) were evaluated at the protein level for expression in human MNV membranes. TIMP3, which showed significant reduction in MNV in all three systems, was abundant in the pathologic basal laminar deposit (BlamD) of the MNV but depleted from Bruch's membrane (BrM) compared to the adjacent choroid (Figs. 6A, 6D). Labeling with antibodies directed against NNMT showed a striking degree of labeling within the MNV compared to the adjacent choroid, with intense reactivity of elongated cells within the areas of subretinal fibrosis (Figs. 6B, 6E).

DISCUSSION

AMD remains a common cause of global blindness. MNV occurs in approximately 5% to 20% of AMD cases^{37,38} and can result in rapid and severe loss of vision. Although anti-VEGF therapies have been widely successful at protecting vision in the setting of MNV, resistance to anti-VEGF therapy is common and can reduce the therapeutic effect, leading to visual deterioration.³⁹ Identifying new molecular features of neovascular choroidal endothelial cells may help advance understanding of MNV triggers and provide insight into treatment resistance to guide new potential therapeutic targets.

Bulk^{40,41} and single-cell^{13,14,17,42,43} transcriptomic experiments have allowed for the study of gene expression in choroidal endothelial cells across health, AMD, and rodent models of neovascularization. Likewise, single-cell RNA sequencing has been utilized to study gene expression changes in models of retinal degeneration.^{44,45} However, MNV, like many other retinal conditions, is a highly regional disease in which focal patterns of pathology are often directly adjacent to more normal retina, RPE, and choroid. It is challenging to use single-cell based approaches to study such heterogenous, spatial disease processes because within each sample, there is no ground-truth classification that labels if cells originate from a healthy versus diseased area. Spatial RNA sequencing addresses this limitation by allowing an expert to label areas of pathology, and this facilitates more specific gene expression comparisons between the lesion and non-diseased areas.

The spatial technology used in this study had a resolution of 55 micrometers. This spot diameter was not near the single-cell level of resolution, and investigation of hematoxylin and eosin images revealed that most spots were covered by several cells and usually represented a mix of different cell types (see Figs. 1, 2). However, this level of resolution still allowed for the investigation of gene expression of different retinal, RPE, and choroidal cell types. For example, known retinal ganglion cell, photoreceptor, and RPE specific genes clearly localized to spots in different retinal layers (see Figs. 2C, 2E, Supplementary Fig. S1). In addition, a deconvolution algorithm estimated the cell type composition of each spot (see Fig. 2), and retinal layers were easily differentiated by their unique deconvoluted compositions.

Capturing the precise spatial location of each spot also allowed for new differential expression analyses. For example, we identified genes within the neural retina that had varied expression along a continuous gradient from the center of each section (and nearest the fovea) to the edges (see Fig. 5). The identified foveal-enriched genes were highly correlated with previous single-cell level investigations, highlighting the power of spatial technology to perform nuanced biological comparisons. As technology evolves, similar analyses in which the spatial location of gene expression is investigated along a distance gradient may provide insight into the regional predilection of other retinal conditions, such as inherited retinal diseases.

The primary aim of this study was to identify genes dysregulated within the MNV. Although we acquired 3 different sections through the MNV lesion, with a diameter of 55 microns, no spot was entirely contained within the carefully traced neovascular membrane. As such, applying the CSIDE algorithm allowed us to control for the two major sources of variation in differential expression: cell type composition and MNV overlap. Instead of setting a cutoff value and arbitrarily classifying which spots were in and out of the neovascular membrane, CSIDE permits continuous explanatory variables. Therefore, we used ImageJ to quantify the intersection between each spot and the carefully traced MNV and used this value to identify genes associated with MNV overlap.

MNV-enriched genes expressed by RPE cells and endothelial cells were predicted to belong to known angiogenesis and neovascular pathways (see Fig. 3). For example, the MNV lesion was enriched in integrins (such as endothelial ITGB1 and RPE ITGB8) previously associated with angiogenesis and blood vessel network formation.^{24,29} Likewise, VEGF and TGFB1, which have been extensively associated with neovascularization, were identified as two of the most likely upstream regulators of the observed MNVenriched genes predicted to originate from endothelial cells and RPE cells (see Figs. 3I, 3J). In addition, we compared MNV-enriched genes identified by the current spatial study with a previous single-cell experiment of two MNV donors¹⁴ as well as a rodent model of choroidal neovascularization.¹³ The most upregulated endothelial MNV gene in the current study, IL6, has been previously shown to be expressed by choroidal macrophages and associated with neovascularization in rodents.³⁵ Collectively, these findings suggest that at the current resolution, spatial RNA sequencing can identify known pathogenic drivers of MNV and offers a useful approach to study gene expression within this highly spatial disease process.

There are several limitations to this study. First, as mentioned above, the 55-micrometer diameter resolution of the spatial technology captured several cells within each spot. We partially addressed this limitation by using the nonparametric differential expression algorithm CSIDE, which deconvoluted cell type composition within each spot and stratified differential expression results by cell type. However, such deconvolutions are imperfect, and the unambiguous determination of cell types would be improved with greater spatial resolution. Second, the MNV donor eye had a greater than 5-year history of neovascular AMD. Although endothelial cells and other components of MNV membranes may be dynamic, with growth and revascularization, the gene expression changes observed within the MNV may reflect late stage or remodeled disease, which may be distinct from expression patterns observed earlier in the disease course. It is possible that earlier, disease-initiating molecular changes are no longer detectable at this time point. However, expression patterns within a longstanding type 1 neovascular membrane still provide insight into homeostasis of this pathology and may shed light on potential mechanisms of neovascularization and anti-VEGF resistance. Third, only one MNV donor and one age-matched control donor were used in the current study. Whereas this MNV donor represents a valuable sample with correlated clinical imaging, histopathology, and spatial gene expression, additional validation of the observed expression changes in other human donors is required.

Spatial gene expression is a rapidly evolving technology.⁴⁶ Recently, commercialized systems allow for true single-cell or subcellular resolution of gene expression within a spatial context (e.g. Refs. 47 and 48). However, such technologies use probe-based hybridization and thus necessitate custom gene panel designs. Similarly, unbiased sequencing-based platforms are improving spot resolution to the near single-cell level. This current investigation demonstrates that spatial technology can be used to query gene expression in focal areas of pathology and along regional distance gradients within the heterogenous retina, RPE, and choroid. Further studies, along with technological advances, will continue to improve our understanding of pathogenesis mechanisms in MNV and other regional retinal diseases. Insights from this potent technology may guide the development of future targeted therapeutic interventions with potential to decrease vision loss from AMD and MNV.

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