Comprehensive analysis of gene expression in human retina and supporting tissues

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Understanding the influence of gene expression on the molecular mechanisms underpinning human phenotypic diversity is fundamental to being able to predict health outcomes and treat disease. We have carried out whole transcriptome expression analysis on a series of eight normal human postmortem eyes by RNA sequencing. Here we present data showing that ∼80% of the transcriptome is expressed in the posterior layers of the eye and that there is significant differential expression not only between the layers of the posterior part of the eye but also between locations of a tissue layer. These differences in expression also extend to alternative splicing and splicing factors. Differentially expressed genes are enriched for genes associated with psychiatric, immune and cardiovascular disorders. Enrichment categories for gene ontology included ion transport, synaptic transmission and visual and sensory perception. Lastly, allele-specific expression was found to be significant for CFH, C3 and CFB, which are known risk genes for age-related macular degeneration. These expression differences should be useful in determining the underlying biology of associations with common diseases of the human retina, retinal pigment epithelium and choroid and in guiding the analysis of the genomic regions involved in the control of normal gene expression.

INTRODUCTION

Genome-wide association studies (GWASs) have successfully identified many single-nucleotide polymorphisms (SNPs) associated with complex diseases. Most of these associated SNPs reside within specific genomic regions that include dozens of nearby genes, falling short of proving causation. This is because ∼88% of the genetic variants currently associated with complex diseases and traits by GWAS lie within intronic or intergenic regions (1). Recent GWASs for eye diseases/traits have delivered a number of novel findings across a diverse range of diseases, including age-related macular degeneration (AMD), glaucoma and refractive error (2–4). However, despite this astonishing rate of success, the major challenge still remains to not only confirm that the genes implicated in these studies are truly the genes conferring protection from or risk of disease but also to define the functional roles these genes play in disease. Current thinking is that these nucleotide variations are likely to have causal effects by influencing gene expression rather than affecting protein function (5). A growing number of studies have shown that these SNPs influence gene expression variation in the population (6,7). Global gene expression studies—not acquiring a priori hypotheses—provide a method to investigate the pathogenesis of common disorders on a large scale (8).

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The neurosensory retina is a specialized neural tissue lining the back of the eye that is responsible for processing images and sending these images to the brain. It originates as an outgrowth of the brain during ontogenesis and is thus part of the central nervous system. The cellular composition of the retina is complex with a variety of cell types. The macula has the highest overall density of neurons and is responsible for central vision. The retinal pigment epithelium and choroidal vasculature constitute the photoreceptor support system. The choroid is part of the systemic circulation, i.e., outside the blood retina barrier, and contains multiple resident and transient cells. The sclera is the protective structure of the eye and encloses the eyeball except in the region of the cornea.

Molecular phenotypes such as gene and protein expression are influenced by nearby DNA variations. Therefore, to uncover the genetic determinants regulating expression in tissues involved in chorioretinal diseases, we profiled eight human neurosensory retina and retinal pigment epithelium (RPE)/choroid/sclera (RCS) samples by RNA sequencing (RNA-Seq). We also genotyped DNA from each sample on an Omni 2.5 million SNP chip. Within the RCS sample, the RPE cells and the choroidal vasculature form the photoreceptor support system and are preferentially affected in AMD.

Previous studies on gene expression of the eye have been limited to microarrays (9–14). Our study is the first to report on the differential expression and differential alternative splicing between the macula and peripheral regions of the retina and RCS using RNA sequencing. Our comprehensive analysis of the retina and RCS gene expression profiles revealed large differences in expression between these layers of the posterior eye and a smaller but significant number of differences between the macula and periphery of each layer. Specifically, we identified 9555 differentially expressed (DE) genes between retina and RCS layers in macula and >10 149 DE genes in periphery. We identified >2051 DE genes between peripheral and macular retina (MR) and 926 between peripheral RCS (PRCS) and macular RCS (MRCS). Our differential alternative splicing (DAS) analysis also revealed differences between layers and location and varied from a low of two DAS events between MRCS and PRCS and a high of 575 DAS events between peripheral retina (PR) and PRCS. Significantly, we replicated a portion of our DE genes by QPCR and DAS events by PCR, demonstrating that our data analysis is reproducible. Finally, we have preliminary evidence demonstrating potential allele-specific expression for CFH, C3 and CFB, three genes that have been previously reported to be associated with AMD (15). These marked differences in gene expression between the macula and periphery of each layer may offer some explanation for why some diseases that primarily affect the macula and others affect the periphery. The marked differences between retina and RCS also underscore the tremendous diversity of biological processes required to maintain good photoreceptor health.

RESULTS
Overall quality parameters of the RNA-Seq dataset

We used RNA-Seq to characterize the chorioretinal transcriptomes in a Discovery set of eight normal human eyes. For each eye, we sequenced four RNA-Seq samples and generated close to 100 million 101-bp paired-end reads per sample. We mapped the sequence reads to the reference human genome (hg19) using GSNAP (16). Our data are of high quality with 76–94% of the reads mapped to the human genome and 60–81% retained after stringent quality control filtering, among which 86–93% mapped to genes defined by RefSeq (Supplementary Material, Table S1).

Analysis of gene expression
Global analysis of gene expression

We first considered the overall gene expression in each sample. Using filtered mapped reads, we estimated the expression levels of 23 569 RefSeq protein-coding genes using the fragments per kilobase of gene per million mapped fragment (FPKM) metric (17). With coverage depth ranging from 66 to 133 million paired-end reads per sample, we detected the expression (i.e., FPKM > 0) of the majority of known protein-coding genes. The average number of expressed genes was 18 318 (78%) in MR, 17 334 (74%) in MRCS, 18 850 (80%) in PR and 19 022 (81%) in PRCS (Table 1). Although the overall numbers of expressed genes were similar in the four groups, there was an excess of moderate-to-low-abundance (FPKM < 5) genes in retina as compared with RCS (MR versus MRCS: Fisher’s exact test \( P = 7.8 \times 10^{-4} \); PR versus PRCS: \( P = 2.0 \times 10^{-3} \)). We also observed a slight excess of moderate-to-low-abundance genes in periphery as compared with macula in the RCS layer \( (P = 0.029) \). There was a similar trend in retina (MR versus PR: \( P = 0.25 \)). However, since these tests were not corrected for multiple testing, the comparison of PRCS versus MRCS would not be statistically significant if multiple comparisons were considered.

Based on the estimated FPKM values in each sample, we then investigated the similarity in the global gene expression profiles among the four tissue groups using average linkage hierarchical clustering of the top 1000 most divergent genes defined by the coefficient of variation of the FPKMs (Fig. 1A). The retina samples clearly separated from the RCS samples, suggesting that contamination of retina and RCS during dissection was negligible. Moreover, there was a clear distinction between macula and periphery for retina, but the distinction between macula and periphery was less clear for RCS. These results suggest the existence of tissue-specific gene expression signatures and that regional differences were less pronounced than differences between tissues.

Table 1. Numbers of expressed genes

| Group   | Minimum | Maximum | Mean  |
|---------|---------|---------|-------|
| MR      | 17 639  | 18 800  | 18 318|
| MRCS    | 16 177  | 18 369  | 17 334|
| PR      | 18 258  | 19 577  | 18 850|
| PRCS    | 18 519  | 20 938  | 19 022|

A gene was considered expressed if the FPKM was >0. The minimum, maximum and mean numbers of expressed genes were calculated based on the eight samples in each eye part.
Differentially expressed genes

To detail the genes expressed in each tissue sample, we carried out differential expression analysis using Cufflinks (18). Figure 1B shows the numbers of genes that were differentially expressed. Consistent with the hierarchical clustering analysis, the difference between tissues (e.g. MR versus MRCS) was larger than the difference between locations within the same tissue layer (e.g. MR versus PR). When comparing retina and RCS, we found 9555 DE genes in macula and 10 149 in periphery. In contrast, a smaller number of DE genes were identified when comparing different locations within the same tissue; for retina, the number of DE genes was 2051, whereas it was 926 for RCS. Among genes that were differentially expressed between macula and periphery, 60–70% had higher expression levels in macula (MR versus PR: \( P < 2.2 \times 10^{-16} \); MRCS versus PRCS: \( P < 2.2 \times 10^{-16} \)). Among genes that were differentially expressed between retina and RCS, \( \sim 52\% \) had higher expression levels in RCS (MR versus MRCS: \( P = 9.2 \times 10^{-4} \); PR versus PRCS: \( P = 2.0 \times 10^{-3} \)). Most of the DE genes (\( \sim 70\% \)) had at least 2-fold gene expression difference, except for the comparison of PR and MR in which only \( \sim 39\% \) had a gene expression difference of >2-fold. These results agree with the patterns shown in scatter plots of the estimated FPKMs (Fig. 1C and D).

As our differential expression analysis results were based on Cufflinks v2.1.1, we decided to utilize additional statistical programs to confirm the results. We analyzed the data using two other programs, edgeR (19) and DESeq (20), which utilize independent statistical methodologies (Supplementary Material, Fig. S1). Focusing on the DE genes identified by Cufflinks v2.1.1, we found agreement among all three programs for 8378 (88%) DE genes, when comparing MR and MRCS, 8929 (88%) for the comparison of PR and PRCS, 1195 (58%) for MR and PR and 203 (22%) for MRCS and PRCS (Supplementary Material, Fig. S1). The relatively lower percentage of agreement in the comparison of MR and PR, and the comparison of...
MRCS and PRCS is due to the fact that many DE genes identified by Cufflinks v2.1.1 had a small fold change (<2) of gene expression, and these subtle changes were possibly missed by edgeR or DESeq owing to their lack of power. This was confirmed by the fact that >99% of these DE genes showed the same direction of gene expression changes across all three methods. If we required agreement with either edgeR or DESeq only, then the percentage of agreement would increase to 84% for the comparison of MR and PR and 64% for MRCS and PRCS. These comparisons suggest that our findings from Cufflinks v2.1.1 are reliable. Hereafter, we will only report results based on Cufflinks v2.1.1.

Enrichment analysis of differentially expressed genes

To gain more insight into biological pathways that possibly explain the differences among different tissue layers of the eye, we carried out functional annotation analysis using Database for Annotation, Visualization and Integrated Discovery (DAVID) (21). For the comparison between MR and MRCS (Table 2; Supplementary Material, Table S2A), DE genes with higher expression levels in MR were significantly enriched as follows: for psychiatric disorders, chemical dependency and neurological disorders in disease category; for synaptic transmission, sensory perception of light stimulus and visual perception in gene ontology (GO); and for long-term potentiation, MAPK signaling pathway and endocytosis in Kyoto Encyclopedia of Genes and Genomes (KEGG). Genes with higher expression levels in MRCS (Supplementary Material, Table S2B) were significantly enriched as follows: for immune, cardiovascular and infection in disease category; for immune system processes, positive regulation of biological process and response to external stimulus in GO; and for ribosomes, cell adhesion molecules and focal adhesion in KEGG. Enrichment analysis for DE genes between PR and PRCS revealed a similar pattern (Table 2; Supplementary Material, Table S2C and 2D).

We also carried out functional annotation analysis for genes that were differentially expressed between macula and periphery (Table 3). For the comparison in retina, the DE genes that had higher expression levels in MR (Supplementary Material,

### Table 2. DAVID enrichment analysis for DE genes between retina and RCS

| Category          | MR > MRCS Term | Gene count | P-value | MR < MRCS Term | Gene count | P-value |
|-------------------|----------------|------------|---------|----------------|------------|---------|
| **Disease class** |                |            |         |                |            |         |
| Psych             | 195            | 2.5 × 10\(^{-19}\) | Immune | 383 | 1.4 × 10\(^{-11}\) |
| Chemdependency    | 57             | 4.3 × 10\(^{-4}\) | Cardiovascular | 328 | 4.4 × 10\(^{-10}\) |
| Neurological      | 140            | 2.5 × 10\(^{-2}\) | Infection | 165 | 7.3 × 10\(^{-10}\) |
| **Gene ontology** |                |            |         |                |            |         |
| Synaptic transmission | 151 | 1.1 × 10\(^{-23}\) | Immune system process | 476 | 2.9 × 10\(^{-45}\) |
| Sensory perception of light stimulus | 119 | 2.1 × 10\(^{-22}\) | Positive regulation of biological process | 802 | 1.1 × 10\(^{-38}\) |
| Visual perception | 119            | 2.1 × 10\(^{-22}\) | Response to external stimulus | 423 | 4.5 × 10\(^{-36}\) |
| **KEGG pathway**  |                |            |         |                |            |         |
| Long-term potentiation | 35        | 1.4 × 10\(^{-8}\) | Ribosome | 79 | 1.5 × 10\(^{-26}\) |
| MAPK signaling pathway | 84      | 9.7 × 10\(^{-5}\) | Cell adhesion molecules (CAMs) | 83 | 2.3 × 10\(^{-10}\) |
| Endocytosis       | 59             | 1.8 × 10\(^{-3}\) | Focal adhesion | 113 | 4.5 × 10\(^{-10}\) |
| **PR > PRCS**     |                |            |         |                |            |         |
| Psych             | 190            | 1.6 × 10\(^{-10}\) | Immune | 392 | 1.9 × 10\(^{-12}\) |
| Chemdependency    | 57             | 6.4 × 10\(^{-4}\) | Infection | 170 | 1.1 × 10\(^{-10}\) |
| Neurological      | 141            | 3.2 × 10\(^{-2}\) | Cardiovascular | 324 | 3.8 × 10\(^{-8}\) |
| **Gene ontology** |                |            |         |                |            |         |
| Cellular process  | 2701           | 7.0 × 10\(^{-26}\) | Immune system process | 478 | 5.6 × 10\(^{-39}\) |
| Synaptic transmission | 154       | 4.9 × 10\(^{-21}\) | Positive regulation of biological process | 819 | 6.1 × 10\(^{-34}\) |
| Sensory perception of light stimulus | 121 | 1.2 × 10\(^{-21}\) | Response to external stimulus | 424 | 1.8 × 10\(^{-30}\) |
| **KEGG pathway**  |                |            |         |                |            |         |
| Oxidative phosphorylation | 54        | 6.3 × 10\(^{-6}\) | Ribosome | 67 | 1.7 × 10\(^{-13}\) |
| Alzheimer’s disease | 61           | 3.1 × 10\(^{-5}\) | Cell adhesion molecules (CAMs) | 81 | 2.1 × 10\(^{-8}\) |
| Parkinson’s disease | 51           | 2.5 × 10\(^{-5}\) | Focal adhesion | 112 | 1.5 × 10\(^{-8}\) |

P-values were Bonferroni corrected. Displayed are the top three enriched terms for each category.

### Table 3. DAVID enrichment analysis for DE genes between macula and periphery

| Category          | MR > PR Term | Gene count | P-value | MR < PR Term | Gene count | P-value |
|-------------------|--------------|------------|---------|--------------|------------|---------|
| **Gene ontology** |              |            |         |              |            |         |
| Ion transport     | 110          | 3.1 × 10\(^{-12}\) | Translational elongation | 110 | 2.6 × 10\(^{-22}\) |
| Metal ion transport | 7          | 3.3 × 10\(^{-11}\) | Sensory perception of light stimulus | 77 | 2.0 × 10\(^{-13}\) |
| Transmission of nerve impulse | 64 | 4.7 × 10\(^{-11}\) | Visual perception | 64 | 2.9 × 10\(^{-13}\) |
| **KEGG pathway**  |              |            |         |              |            |         |
| No significant enrichment | MRCS > PRCS | MRCs < PRCS |
| System process    | 167          | 3.2 × 10\(^{-38}\) | Cell adhesion | 32 | 1.9 × 10\(^{-4}\) |
| Muscle contraction | 51          | 4.5 × 10\(^{-24}\) | Biological adhesion | 28 | 1.0 × 10\(^{-4}\) |
| Multicellular organimal process | 289 | 5.2 × 10\(^{-31}\) | Cell development | 28 | 1.3 × 10\(^{-3}\) |
| **KEGG pathway**  |              |            |         |              |            |         |
| Dilated cardiomyopathy | 21         | 2.9 × 10\(^{-7}\) | TGF-beta signaling pathway | 9 | 1.9 × 10\(^{-2}\) |
| Hypertrophic cardiomyopathy | 19 | 1.9 × 10\(^{-7}\) |              |            |         |
| Calcium signaling pathway | 23 | 4.9 × 10\(^{-5}\) |              |            |         |

P-values were Bonferroni corrected. Displayed are the top three enriched terms for each category.
ASE ratio was calculated as the fraction of the alternative allele among all alleles at the SNP. All three SNPs were associated with AMD.

### Table 4. Evidence of ASE at GWAS-associated SNPs for eye diseases

| SNP      | Position   | Gene | Reference allele | Alternative allele | Subject | Sample | ASE ratio | P-value |
|----------|------------|------|------------------|--------------------|---------|--------|-----------|---------|
| rs2230199| chr19:6718387 | C3   | G                | C                  | 11–1516 | PR     | 0.75      | 2.5E-02 |
|          |            |      |                  |                    | 11–1516 | PRCS   | 0.73      | 3.6E-09 |
|          |            |      |                  |                    | 11–1516 | MRCS   | 0.65      | 3.75E-05|
|          |            |      |                  |                    | 11–2043 | MR     | 0.60      | 6.33E-05|
| rs1061147| chr1:196654524 | CFH  | A                | C                  | 11–1516 | PRCS   | 0.46      | 4.42E-02|
|          |            |      |                  |                    | 11–1516 | PR     | 0.47      | 4.77E-02|
|          |            |      |                  |                    | 11–1624 | MRCS   | 0.41      | 6.89E-08|
|          |            |      |                  |                    | 11–1614 | MRCS   | 0.69      | 4.11E-02|
| rs641153 | chr6:31914180 | CFB  | G                | A                  | 11–1833 | PRCS   | 0.66      | 2.93E-04|

ASE ratio was calculated as the fraction of the alternative allele among all alleles at the SNP. All three SNPs were associated with AMD.
Biological replication of differential expression

A total of 204 genes detected to be DE by Cufflinks from the Discovery set were chosen for biological replication based on their biological relevance and significance of the P-values. The Replication set included eight normal eyes, but two MR samples were eliminated owing to poor RNA quality. Paired t-test was used to determine P-value for differential expression. Percentages of DE genes replicated were as follows: MR versus MRCS (61% replicated), PR versus PRCS (83% replicated), MR versus PR (59% replicated) and MRCS versus PRCS (29% replicated).

Analysis of alternative splicing

Global analysis of alternative splicing

RNA-Seq’s specificity for closely related transcript isoforms allowed us to go beyond gene expression level and identify isoform differences owing to pre-mRNA alternative splicing (AS). Using MATS (28), we identified a large number of AS events, particularly in retina (Fig. 2A). The average numbers of AS events were 15 560 in MR and 15 290 in PR, whereas the corresponding numbers were 11 760 in MRCS and 10 050 in PRCS, respectively. These results suggest that many genes were alternatively spliced and the AS patterns were different between the retina and RCS tissue layers.

For insight into AS patterns in different chorioretinal tissues and locations, we carried out hierarchical clustering analysis based on the top 150 most divergent AS events defined by the coefficient of variation of the exon-inclusion levels (Fig. 2B). Interestingly, the retina samples clearly separated from the RCS samples, although only 150 AS events were used in the clustering. However, within each tissue, the distinction between macula and periphery was less clear with retina showing a slightly better separation of macula and periphery. These results suggest the existence of tissue-specific AS signatures, but the difference between locations within the tissue layer was subtle.

Differential alternative splicing

Next, we examined whether an AS event showed different degrees of splicing between different chorioretinal tissues of the eye by testing whether the difference in exon-inclusion levels exceeded 5% for any cassette exon of a gene. Figure 2C shows the numbers of differential AS (DAS) events. When comparing retina and RCS, we found 431 DAS events in macula and 575 in periphery. In contrast, we detected a much smaller number of DAS events between MR and PR (39 events) and between MRCS and PRCS (2 events). These results were consistent with the patterns revealed by the hierarchical clustering analysis.

For DAS events identified from the comparison of retina and RCS, we further examined their exon-inclusion levels (Fig. 2D and E). In macula, more DAS events appeared to have higher exon-inclusion levels in retina than in RCS (MR versus MRCS: P = 0.027). There was a similar trend in periphery, but the difference was not significant (PR versus PRCS: P = 0.13).

We recognize that the analysis of DAS is challenging because detection of AS events relies on the number of junction reads. To assess the reliability of our findings, we classified DAS events into three tiers according to the degrees of confidence. Tier 1 includes high-confidence events that have been annotated by RefSeq, i.e. both the inclusion and the exclusion isoforms are present in RefSeq’s annotation. Tier 2 includes moderate-confidence events for which the corresponding genes are alternatively spliced according to UniProt. Tier 3 includes the remaining events. Based on these classifications, we confirmed that >80% of the DAS events were in tier 1 for comparisons between MR and MRCS, MR and PR and PR and PRCS. Of the two DAS events identified for the comparison between MRCS and PRCS, one was in tier 1 and the other was in tier 2 (Fig. 2F). These results suggest that our findings on DAS were reliable.

Splicing factors are RNA-binding proteins that play key roles in coordination of spliceosome functions and regulation of AS. We compiled a list of 60 well-studied splicing factors and examined their expression in all samples. Thirty-seven splicing factor genes were expressed in chorioretinal tissues: 12 being DE between MR and MRCS, 16 between PR and PRCS and 11 DE between retina and RCS in both regions. As expected, the number of DE genes was much smaller when comparing different regions; only three genes were differentially expressed between MR and PR, and none were differentially expressed between MRCS and PRCS. These findings are consistent with the number of DAS events in that the number of DAS events identified in each comparison perfectly correlated with the number of splicing factors that were differentially expressed, suggesting that the differences in DAS events were likely due to the expression difference of related splicing factors.

Enrichment analysis of genes with differential alternative splicing

Next, we carried out functional annotation analysis for genes showing DAS when comparing retina and RCS. For both macula and periphery, genes were significantly enriched for actin filament-based process, cytoskeleton organization and anatomical structure development in GO, and for tight junction in KEGG. We did not perform functional annotation analysis for the comparison between macular and periphery owing to the small number of genes showing DAS.

Overlap with GWAS signals on eye diseases

For genes showing DAS, we further examined whether they had been implicated in recent GWAS on AMD (15), glaucoma (29) and myopia (30) based on the NHGRI GWAS catalog. Among the 91 unique GWAS genes we examined, three genes—CD46, DDR1 and RREB1—showed DAS between PR and PRCS. Interestingly, all three were identified from GWAS on AMD (15), and all had higher expression levels in RCS than in retina. Of these genes, CD46 (also called membrane cofactor protein or MCP) is notably the only membrane-bound regulator of complement that is expressed on the basolateral RPE (22,31). DDR1 is a hypoxia-induced gene involved regulation of cell growth and is expressed in lamina cribrosa cells in the optic nerve (32). We did not find overlap with GWAS signals for genes showing DAS identified from other comparisons.

Biological replication of differential alternative splicing

Five genes showing DAS by MATS were chosen for replication in a Replication set of an additional eight normal postmortem eyes. All five genes were replicated in all eight eyes demonstrating the robustness of our analysis. Figure 3 shows gel images for two representative eyes.
DISCUSSION

GWASs have been used to correlate genetic variants with disease in a hypothesis-generating manner. Linking these genetic variants with causation has been the next major milestone, with tissue specificity, low resolution of DNA genotypes and technical challenges of assaying molecular traits being the limiting factors to date. Previous studies have examined ocular...
gene expression by RT-PCR and microarrays but have not yet attempted the entire transcriptome of human neurosensory retina and photoreceptor support system by RNA-Seq. In the present study, we conducted a comprehensive RNA-Seq analysis of MR and PR and macular and PRCS from eight phenotypically well-characterized normal adult eyes. The molecular signatures that we obtained for macula and periphery will be very useful biomarkers for characterizing different stages of chorioretinal disease. Our results for RNA-Seq are to our knowledge novel owing to the comparison of MR and PR and for the same regional comparison of the RCS.

RNA-Seq findings from multicellular tissues should be interpreted in light of the cellular composition of the tissue samples. Human neurosensory retina is dominated by densely packed neurons and includes significant minority populations of radial glia (Müller) cells, astrocytes, microglia and vascular endothelial cells and pericytes. RCS samples, in contrast, include multifunctional RPE, multicellular choroid (with vascular and lymphatic endothelia, fibroblasts, melanocytes, mast cells, autonomic neuronal ganglia and resident and transient cells of monocyte lineage) and structural sclera (with sparse fibroblasts and myofibroblasts) (34,35). To effectively study and understand biological phenomena at the molecular level, variability in gene expression must be taken into account. This variability in gene expression can exist for two reasons. First, the tissue may be dominated by a large number of different cell populations in differing amounts, which can result in a dilution of the signal from less-populated cells below the lower detection limit for RNA-Seq. Second, cell populations can average out individual cellular co-expression patterns, making it difficult to

Figure 3. RT-PCR biological replication of differential alternative splicing (DAS) events. (A) Replication of TIMM8B1 DAS event from PR versus PRCS comparison. (B) Replication of STXB1 DAS event from PR versus PRCS comparison. (C) Replication of CSDA (YBX3) DAS event from PR versus PRCS comparison. (D) Replication of PKP4 DAS event from PR versus PRCS comparison. (E) Replication of FMNL1 DAS event from MR versus MRCS comparison.
determine whether two RNAs always occur within the same cell. We hypothesized that our expression from the retina was not sensitive enough to detect gene expression from less abundant retinal cells. We expected neurosensory retina RNA expression to be dominated by the transcriptome of the numerically dominant photoreceptor population (>100 M cells), yet we found robust RNA-Seq signal for genes expressed by less populous neuronal and glial cell classes (36). The cell classes include POU4F2 (ganglion cells) (37), SNAP25 (horizontal cells) (38), CALB2 (amacrine cells) (39), RCVRN (bipolar cells) (39) and GLUL (Müller cells) (40). FPKM values for all five genes varied from a low of 1.0 for POU4F2 to a high of 4148 for RCVRN, suggesting that we had excellent representation of the retina’s less numerous cells. Next, we asked whether signal for specific neuronal types was also represented in the RNA-Seq data. We found high RNA expression for CBP5 (39) (specific for bipolar cells DB3, DB4, RB; 258 FPKM) and SYT2 (39) (specific for amacrine cells A8; 23 FPKM) among others, suggesting that our utilization of whole retina RNA was sensitive enough to detect RNA expression from less common retinal cells. We did not interrogate the RCS samples with the same questions because cell populations in those tissues are not as well defined quantitatively as they are in neurosensory retina, yet we would expect on the basis of this analysis that all RCS cell types have been adequately represented as well.

The largest difference in expression in these eight unremarkable adult human eyes occurs between the neurosensory retina and the RCS. In total, 9555 genes in macula, and 10 149 genes in periphery were differentially expressed between layers, and these were evenly distributed between those that were higher in retina and those that were higher in RCS in both locations. Of the genes enriched in the retina as compared with RCS and those enriched in MR as compared with PR, a number of genes were found to be associated with psychiatric disorders following DAVID analysis (41). We examined the top 25 most significant DE genes for each pairwise comparison (MR versus MRCS, PR versus PRCS and MR versus PR). This yielded a list of 50 unique genes. Following an expansive literature review, we found that 45 of them were implicated in one or more psychiatric disorders, including schizophrenia (n = 24), Alzheimer’s disease (n = 8), autism (n = 8), depression (n = 6) and bipolar disorder (n = 6) (Supplementary Material, Table S5). Of these 50 genes, mutations in eight are known to have a specific eye phenotype in humans. These include DDC, GAD1, GAD2, MAPT, PAX6, PPP2R2B, SYP and TFAP2B. Common eye phenotypes seen include oculomotor apraxia, ptosis, nystagmus and opthalmoplegia. One gene of particular interest is AQP4, aquaporin 4. Enhanced AQP4 expression has been found in the brains of patients with Alzheimer’s disease (42). An eye phenotype in humans has not been documented, but Aqp4tm1Ask/Aqp4tm1Ask mice display significantly reduced electroretinogram (ERG) b-wave potentials (43). Patients with Alzheimer’s disease are known to display attenuated pattern- and multifocal-electroretinogram responses as well as decreases in retinal nerve fiber layer thickness as determined by optical coherence tomography (OCT) (44–47). Together, these findings suggest that ERG b-wave response and OCT may be useful tools to identify individuals at risk for developing Alzheimer’s disease.

We further explored tissue-specific variation in chorioretinal gene expression by examining the top 25 most significant DE genes between retina and RCS for both macula and periphery (Supplementary Material, Table S6). Many DE genes preferentially expressed in RCS fall into the cardiometabolic category. They are a diverse group, expressing in muscle, vessel walls and blood cells, and they are well explored in systemic vascular diseases. The top genes expressed at higher levels in RCS than retina in both macula and periphery include BDKRB2, CYP4B1, DES, HBB, JPH2 and SAA1. Of these, BDKRB2, DES and HBB have been investigated with regard to the retinal circulation in the context of diabetic retinopathy and remain to be explored in choroid. SAA1 encodes an apolipoprotein and acute phase responder found in isolated RPE-capped drusen (48). The top genes expressed at higher levels in RCS than retina for macula only (Supplementary Material, Table S6) notably include COL1A1, COL1A2 and COL3A1, which encode structural collagens, consistent with the fact that choroid and sclera are both at their thickest under the macula and thus contain many fibroblasts and myofibroblasts (27,49). Polymorphisms of COL genes are associated with systemic hypermobility of joints and disorders of vascular disintegration. Other highly expressed macula RCS genes of interest for roles in AMD initiation and/or progression are CR1 (found in RPE, BrM and blood leukocytes), IL6 (an acute mediator of inflammatory responses found in aqueous of AMD eye), TIBS1 (an antiangiostatic protein reduced in AMD BrM) and VCAM1. The top genes expressed at higher levels in RCS than retina in periphery only include many cellular receptors (ADRA1A, ADRA2C, AGTR1, CD36, NPR1 and NR1H4). RCS expression of these genes is apparently unmasked by thinner neurosensory retina in this region. As the support system for the photoreceptors, the RCS exhibits diverse gene expression in service of maintaining the health of these cells for good visual function. An important next step will be assigning these expression patterns to the multiple RCS cell types.

As an external validation of genes important in cardiovascular disease, we compared our chorioretinal RNA-Seq expression data to microarray gene expression data for chorioretinal cholesterol pathways published by Zheng et al. (50), who analyzed neurosensory retina and RPE of six human donor eyes using a commercially available microarray. This comparison is of interest because esterified cholesterol and phosphatidylcholine are the major volumetric components of drusen (48). AMD’s pathognomonic extracellular lesions located on the inner surface of the choroid, and unesterified cholesterol is present in subretinal drusenoid deposit (reticular pseudodrusen), a lesion located between the photoreceptors and the RPE in many AMD eyes (51,52). Zheng et al. (50) reported expression levels for 84 genes in these two tissues normalized against 6 housekeeping genes. For these 84 genes, we ranked expression levels within our MR, MRCS, PR and PRCS groups, from highest to lowest and compared their ranks to the Zheng dataset (50) (Supplementary Material, Fig. S2). Two observations can be made. First, all microarray genes were also detected by RNA-Seq in all four tissue sample groups. Second, RNA-Seq expression levels were strongly positively correlated with microarray results for these genes. We note that agreement was higher for MR and PR (0.67) than for MRCS and PRCS (0.51–0.54). These findings are plausible because both studies assayed retina, whereas Zheng et al. (50) assayed RPE only versus RCS for RNA-Seq. We can conclude that many major genes in cholesterol and lipoprotein
pathways are expressed and likely functional in retina, RPE and choroid. This comparison comprised an external validation of our RNA-Seq results with regard to a focused set of genes. These corroborative findings also emphasize that pathogenic models based on GWAS-identified SNPs in genes well known from plasma HDL metabolism and expressed in liver, intestine and adipose tissue should incorporate the intraocular expression of these genes.

Another striking finding is that >2000 genes were differentially expressed between MR and PR and >900 genes between MRCS and PRCS. These genes were slightly tilted toward those that were more highly expressed in RCS versus retina. The differential expression between macula and periphery is attributable in part to differences in the cellular composition of these regions. For retina, rod photoreceptors are the most abundant cell type (100 million) (53) with cones more numerous in macula than they are in the periphery but still a minority (<10% of total in macula). The cone-dominated fovea, <0.8 mm in diameter, contains 32 000 cones (53,54). The ratio of macula photoreceptor densities to peripheral photoreceptor densities is 1.9 for cones (because cones dominate a small area) and 2.7 for rods (53,55). A feature unique to macula is the Henle fiber layer, which contains processes of cone and rod photoreceptors transmitting graded potentials from the light-capturing inner segment to synaptic terminals, interleaved by parallel processes of Müller cells. Individual Henle fibers can be up to 600 μm long, and collectively the Henle fiber component of the anatomical outer plexiform layer (Fig. 4) occupies 14% of macular volume at its thickest portion (56–58). The macula contains half (~750 000) of all brain-projecting ganglion cells of the human retina (59); these cells are dominated by midget cells required for high-acuity vision (60). Mean ganglion cell density is 100-fold greater in macula than in periphery. For the choroid, differential cellular composition may be attributable to blood cells retained within the more highly vascularized macula choroid (61) and numerous melanocytes that impart to the macula its greater pigmentation relative to periphery in this mature age group (62). Therefore, our differential expression results between macula and periphery are expected based on the anatomical differences between these regions and will likely be borne out by regional differences in choroid that are being explored anew through high-resolution imaging.

In conclusion, the transcriptome data presented here is the first report showing differential expression between regions within the retina and RCS by RNA-Seq. The magnitude of these differences is striking and expands our catalog of the gene expression variation that exists in the normal eye. The data underscore the need to not only fully understand the expression differences between layers but also between regions of a layer. Overall biological interpretation of regulatory effects—much like in the case of complex traits—is tissue- and region-dependent, highlighting the value of multiple tissue expression datasets. Gene expression is a critical phenotype that reveals biological properties and allows us to probe cellular functions. Combining statistical methods with relevant sample collections of tissues and cell types from well-phenotyped individuals enables the integrated treatment of biological and epidemiological information in an iterative way. This provides us with the highest possible resolution and will reveal the real causes for disease predisposition.

**Figure 4.** *Ex vivo* imaging and histology of eye contralateral to that used for RNA-Seq. (A and B) Color image of excised preserved macula (A) and fovea (B) from a 79-year-old Caucasian male. Optic nerve is at the left edge of the punch. (C) *Ex vivo* spectral domain optical coherence tomography of excised preserved macula, with neurosensory retina at the top and RPE–choroid–sclera at the bottom. The optic nerve is at the left, and the fovea is the dip in the middle. This fovea has postmortem cystic change (arrowheads). The macula is otherwise unremarkable. (D) 0.8-μm-thick epoxy section of tissue post-fixed with the OTAP method, stained with toluidine blue and scanned. Detachment of the neurosensory retina from the RPE is common in postmortem specimens, even if microscopically well-preserved like this one. Green frame indicates area shown at higher magnification in E. (E) Chorioretinal layers on the foveal slope. This area of the retina is cone-dominated, as evidence by the few darkly stained rod nuclei that contrast with numerous cone nuclei in the ONL. Layers: GCL, ganglion cell layer; IPL, inner plexiform layer; INL, inner nuclear layer; OPL, outer plexiform layer; HFL, Henle fiber layer; IS, inner segments of photoreceptors; OS, outer segments of photoreceptors; RPE, retinal pigment epithelium; Ch, choroid; Sc, sclera. Detached retina was digitally reapposed to RPE for illustrative purposes (Eye 2011001R is shown as a left eye by UAB lab convention. As AEB 11-1515-P, it is contralateral to 11-1516-P analyzed by RNA-Seq.).

This view has the potential to also lead to higher confidence candidates in the absence of direct functional support for any one gene, as might be the case for inconclusive GWAS findings where the SNPs identified have no known functional role. Generating even larger-scale expression datasets may provide a path to more rapidly elucidating not only the genetic basis of eye disease but also the impact of gene expression on molecular networks that in turn induce variations in disease associated traits.

**MATERIALS AND METHODS**

**Eye collection**

This study conformed to Institutional Review Board regulations for use of human tissues at University of Alabama (UAB) and at
University of Pennsylvania (Penn). Our study utilized two sets of eight pairs of eyes from non-diabetic Caucasian donors at a death-to-preservation interval of <6 h. This interval was chosen because RPE RNA quality is adequate during this time frame and begins to decline thereafter (63). A Discovery set of eyes consisted of five males and three females [73.9 year ± 12.5 year (mean ± standard deviation)]. A Replication set also consisted of five males and three females (85.9 ± 3.0 year) with somewhat older ages owing to an adjustment of the tissue collection protocol to maximize the number of AMD eyes collected for separately reported studies. Ocular health histories were not available. Eyes were opened anteriorly by Alabama Eye Bank recovery personnel using an 18-mm-diameter trephine, followed by a radially oriented snip to the iris margin to facilitate penetration of preservatives into the fundus. Preservatives used were RNAlater (Qiagen, Valencia, CA, USA) for the left eye and 2% glutaraldehyde and 1% paraformaldehyde in 0.1 M phosphate buffer for the right eye, both at 4°C. Left eyes were shipped on wet ice via overnight courier to Penn where they were processed upon arrival.

Maculopathy status of right eyes was assessed at UAB by a three-component protocol. Eyes underwent multimodal ex vivo imaging of excised 8-mm-diameter macular punches using digital color photography and spectral domain optical coherence tomography volume scans (SD-OCT; Spectralis, Heidelberg Engineering) with a custom tissue holder. They underwent internal globe examination using a dissecting scope (Nikon SMZ-U) with oblique trans- and epi-illumination in consultation with an MD medical retina specialist (J.A.K.). Finally, eyes were submitted for histopathology using macula-wide high-resolution sections. Macular punches including retina, RPE, choroid and sclera were post-fixed in osmium tannic acid for 6 h. This interval was found to facilitate penetration of preservatives into the fundus. Preservatives used were RNAlater (Qiagen, Valencia, CA, USA) for the left eye and 2% glutaraldehyde and 1% paraformaldehyde in 0.1 M phosphate buffer for the right eye, both at 4°C. Left eyes were shipped on wet ice via overnight courier to Penn where they were processed upon arrival.

Alignment of RNA-Seq reads

The RNA-Seq data were aligned to the hg19 reference genome using GSNAP (16) with default options. In order to eliminate mapping errors and reduce potential mapping ambiguity owing to homologous sequences, several filtering steps were applied. Specifically, we required the mapping quality score of ≥30 for each read, reads from the same pair were mapped to the same chromosome with expected orientations and the mapping distance between members of the read pair was <500 000 bp. Quality control analysis of the aligned data was performed using program RNA-SeQC (33). All subsequent analyses were based on filtered alignment files.

Analysis of differential gene expression

Transcripts were assembled using Cufflinks software (18). For each gene, we compared the expression levels between chorioretinal samples, including MR versus MRCS, PR versus PRCS, PR versus MR and PRCS versus MRCS. The first two comparisons allow us to identify genes that are differentially expressed between different tissues, whereas the last two comparisons allow us to identify genes that are differentially expressed between different locations within the same tissue layer. A gene was considered differentially expressed if the false discovery rate (FDR)-adjusted P-value is <0.05.
Analysis of differential alternative splicing

The availability of RNA-Seq data on the same subject in different tissues and locations provides an opportunity to identify differential alternative splicing (AS) events. To identify such events, we used MATS (28). This software implements a Bayesian approach that detects differential AS under two conditions by examining whether the difference in the exon-inclusion levels between two samples exceeds a given user-defined threshold (0.05 in our analysis). An AS event was declared if the inclusion level of an exon is between 0 and 1. A differential AS (DAS) event was declared if the FDR-adjusted \( P \)-value is < 0.05.

Functional annotation

To identify overrepresented functional categories among genes that are differentially expressed or differentially spliced, we carried out annotation analysis using the DAVID (41). Differentially expressed genes or genes showing DAS were used as input gene list. We looked for enrichment for genetic association with disease class, biological processes in GO and KEGG pathways. Multiple testing was adjusted using the Benjamini–Hochberg approach, and enrichment was declared if the adjusted \( P \)-value is < 0.05.

Biological replication of differential gene expression results via Nanostring

Using the Nanostring nCounter Analysis System (Nanostring Technologies, Seattle, WA, USA), gene expression analysis was conducted on six new phenotypic normal maculas and eight new phenotypic normal peripheral tissues using a custom-designed code set containing 204 genes as previously described (68). These genes were chosen based on biological relevance and having evidence of DE based on Cufflinks analysis (fold change > 2 and an FDR-adjusted \( P < 0.05 \)). Each reaction contained 100 ng of total RNA in a 5 ul aliquot, plus reporter and capture probes. Analysis and normalization of the raw Nanostring data was conducted using nSolver Analysis Software v1.1 (Nanostring Technologies). Raw counts were normalized to internal levels of 14 reference genes: ADAM10, APP, APH1A, FHTT, SNCA, GSTP1, NFE212, DIRAS2, SIRT2, CYC1, GIGYF2, CTIF, YWHAZ and RPL22. The background was subtracted from the data using the average of the negative controls (included in each Nanostring Codeset). Paired t-test was used to assess differential expression. A gene was considered replicated if the \( P \)-value is < 0.05.

Biological replication of differential alternative splicing results via PCR

Replication of our DAS for five genes, TIMM8B1, CSDA, PKP4, STXBP1 and FMNL1, was performed in a Replication set of eight additional eyes. To simplify the replication analysis, genes were chosen from the MATS results that had only two isoforms. Using the genomic coordinates included in the MATS output, the UCSC genome browser was used to visually localize the exon’s location in relation to the flanking exons shared by both isoforms. A portion of the sequence from the exons flanking the differentially spliced exon was taken from sequence data at NCBI and used to create the primers using Primer3 (http://bioinfo.ut.ee/primer3-0.4.0/). To ensure that these primers were target-specific, Primer-BLAST was used (http://www.ncbi.nlm.nih.gov/tools/primer-blast/). Primers are listed in Supplementary Material, Table S7. RNA from the eight replication eyes was extracted using the AllPrep DNA/RNA Mini prep Kit (Qiagen). Extracted RNA samples underwent quality control assessment using R6K ScreenTape on a 2200 Tape station (Agilent) and quantified using Qubit 2.0 Fluorometer (Life Technologies, Foster City, CA, USA). In total, 20 ng of RNA was used to make cDNA using random hexamers for priming and Superscript III first strand synthesis system (Life Technologies). The cDNA was prepared according to the manufacturer’s protocol. The cDNA was amplified using GoTaq Green Master Mix (Promega, Madison, WI, USA) and gene-specific primers. A touchdown PCR method with 68–55 °C temperature gradient was used to amplify the different transcripts. The PCR products were visualized on an Agarose 1.5% gel using ethidium bromide dye.

DNA genotyping and quality control

DNA for the eight Discovery eyes was extracted using AllPrep DNA/RNA Mini Kit. Simultaneous purification of genomic DNA and total RNA from same sample was achieved using this kit. Lysate from each tissue was passed through an AllPrep DNA spin column to selectively isolate DNA and then through RNaseq MinElute spin column to selectively isolate RNA. The extracted genomic DNA quality and quantity was measured using NanoDrop 8000 (Thermo Fisher, DE). All eight DNA samples had a 260/280 ratio of ≥ 1.7. All eyes were genotyped at the Center for Inherited Disease Research using the Illumina HumanOmni2.5-4v1_B SNP array. We performed quality control analysis to ensure the quality of the DNA genotypes. First, we required all samples to have genotype call rate of > 97.5% and no excess or deficient heterozygosity (inbreeding coefficient | F | < 0.1). Next, we performed quality control measures to exclude unreliable SNPs. We eliminated SNPs with genotype call rate of < 98% or if there was significant departure from Hardy–Weinberg equilibrium (\( P < 0.0001 \)).

Analysis of allele-specific expression

To test for allele-specific expression (ASE), we considered exonic SNPs reaching GWAS significance (\( P < 5 \times 10^{-8} \)) for eye-related diseases based on NHGRI GWAS catalog (http://www.genome.gov/26525384). We identified subjects that are heterozygous at the test SNPs. We counted the number of RNA-Seq reads carrying each allele and tested ASE using a one degree of freedom \( \chi^2 \) goodness-of-fit test for equal frequencies of both alleles.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

Conflict of Interest statement. None declared.

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