**Pro-fibrotic pathway activation in trabecular meshwork and lamina cribrosa is the main driving force of glaucoma**

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**ABSTRACT**

While primary open-angle glaucoma (POAG) is a leading cause of blindness worldwide, it still does not have a clear mechanism that can explain all clinical cases of the disease. Elevated IOP is associated with increased accumulation of extracellular matrix (ECM) proteins in the trabecular meshwork (TM) that prevents normal outflow of aqueous humor (AH) and has damaging effects on the fine mesh-like lamina cribrosa (LC) through which the optic nerve fibers pass. Applying a pathway analysis algorithm, we discovered that an elevated level of TGFβ observed in glaucoma-affected tissues could lead to pro-fibrotic pathway activation in TM and in LC. In turn, activated pro-fibrotic pathways lead to ECM remodeling in TM and LC, making TM less efficient in AH drainage and making LC more susceptible to damage from elevated IOP via ECM transformation in LC. We propose pathway targets for potential therapeutic interventions to delay or avoid fibrosis initiation in TM and LC tissues.

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profiles from human TM or LC cultured cells and from human samples of TM or LC with or without POAG, derived from well annotated publicly available datasets (GSE4316, GSE27276, GSE45570, GSE13534, GSE758, E-MEXP-3440, GSE2378, GSE2705). The intracellular SPA analysis is a universal method, which may be used to analyze any physiological, stress, malignancy or other perturbed conditions at the molecular level. In contrast to other existing techniques for aggregation and generalization of the gene expression data for individual samples, our method distinguishes the positive/activator and negative/repressor role of every gene product in each of the distinct signaling pathways analyzed and determines its pathway activation score (PAS).16 Applying our pathway analysis algorithm, we explored signaling pathway activation profiles as a result of pathological conditions and we have discovered that elevated levels of TGFβ in glaucoma-affected tissues are strongly associated with the activation of pro-fibrotic pathways in TM and LC. The elevation in fibrosis-associated signaling may lead to enhanced ECM remodeling, making TM less efficient in humor drainage and causing LC to be more susceptible to damage from elevated IOP via increased tissue stiffness and ECM crosslinking.

Although our findings are exploratory, and are therefore useful primarily for the generation of new hypotheses, given the universal applicability of our platform as a potent in-silico drug screening and efficacy prediction tool,16,17,19 our work may provide a roadmap for potential therapeutic interventions to delay or avoid fibrosis initiation in TM and LC tissues.

Results

**Differential signaling pathway activation in trabecula meshwork tissue of POAG donors**

In order to discover the signaling pathway profiles associated with the onset of glaucoma, glaucoma-related symptoms and secondary effects due to progression of the disease, we performed an in silico pathway activation analysis based on the publicly available datasets (obtained from NCBI GEO and ArrayExpress repositories) on gene-expression studies involving TM and LC. To study the general pathway activation drift in TM of the POAG donors, we applied our pathway analysis algorithm on the transcriptomic data obtained from all known POAG TM datasets, namely, GSE27276 (13 controls and 15 POAG cases) and GSE4316 (3 controls and 2 POAG cases).

The GSE4316 data set contained only 2 POAG samples, making it impossible to estimate the statistical significance of the obtained results. For this reason, the 50 most dysregulated pathways in the TM of these 2 POAG samples compared to controls have been selected based on their corresponding PAS values (Fig. 1A). Since dataset GSE27276 contained a sufficient number of POAG samples to estimate the p-value for each dysregulated pathway, the top 50 differentially activated signaling pathways have been selected based on their statistical significance (p < 0.00002), instead of their corresponding PAS values (Fig. 1B). Despite different criteria used for pathway selection, a substantial overlap in most dysregulated pathways has been seen between the 2 data sets analyzed. Notably, all 50 most dysregulated pathways from the GSE4316 dataset, had also been discovered as differentially activated compared to controls in the analysis of the GSE27276 data set, with p-value <0.05. Interestingly, multiple pro-survival pathways associated with fibrogenesis in different human organs (such as AKT, PAK, p38, ERK, JNK, CREB, cAMP and JAK-STAT) were detected as unregulated in POAG tissue from both datasets (Fig. 1A and B).

**Signaling pathways differentially activated in lamina cribrosa of the POAG donors**

To analyze pathway activation drift as a result of pathophysiological changes in lamina cribrosa during glaucoma progression, we analyzed data set GSE45570 containing samples of optic nerve head (ONH) which includes lamina cribrosa tissues (mixture of ECM, astrocytes, neurons fibers and glial fibrillary acid protein (GFAP) negative lamina cribrosa cells) and dataset GSE13534 containing purified GFAP LC cells respectively from POAG donors and normal control. Given the difficulty of obtaining gene expression data of well annotated, clinically-relevant LC specimens, these were the only POAG data sets currently available. For visualization of pathway activation drift in glaucoma-affected LC, we selected the 50 most significant pathways from the GSE45570 dataset with p-value cutoff 0.05 and the 50 most (judged by PAS value) dysregulated pathways from the GSE13534 data set, since it contains only 2 samples. We found that pathways like JNK, JAK-STAT, PAK, ERK, AKT were up-regulated in POAG ONH (Fig. 1C) whereas ILK, RAS, ERK, PTEN pathways related to adhesion, migration, angiogenesis and Ca2+ signaling were upregulated in POAG LC cells (Fig. 1D). The most significant down-regulated pathways in LC cells were those including the super pathway of cholesterol biosynthesis, PTEN, AKT, Integrin-linked kinase (ILK), Hedgehog and WNT signaling (Fig. 1C and D).

**Effect of increased pressure on signaling pathway activation in cultured ONH astrocytes**

To further elucidate the signaling changes induced by elevated pressure in LC tissue, we have analyzed a transcriptomes derived from cultured ONH astrocytes exposed to a transient increase in hydrostatic pressure (HP), dataset GSE758. Cultured astrocytes were exposed to hydrostatic pressure (60 mm Hg), compared to control astrocytes at ambient pressure (AP), over 6 hr, 24 hr and 48 hr intervals. Our analysis showed that the PAS values of 58, 21 and 40 distinct pathways, were significantly dysregulated compared to controls at 6, 24 and 48 hours respectively (Table S1). To restrict output analysis to pathways that most probably have been dysregulated as a result of increased pressure, we have selected signaling axes that were significantly up- or downregulated at all 3 time intervals. Pathways like the Wnt main pathway, Wnt-PKC, Wnt/CREB3, p38, ILK related to cytoskeletal adhesion complexes, growth hormone and glucocorticoid receptor (inflammatory cytokines) were significantly altered in ONH astrocytes under raised HP conditions, when compared to control cells (Fig. 2). The ILK pathway related to cytoskeletal adhesion was down-regulated 3-fold after 6 hr of exposure, and then up-regulated after 24 hr
Finally, down-regulated after 48 hours (Fig. 2). Although ILK signaling was not previously shown in glaucoma, elevated ILK signaling was associated with increased EMC remodeling in response to mechanical stiffness in lung.20 Therefore, during raised IOP conditions in glaucoma, increased ILK levels may take part in microtubule and microfilament re-arrangement upon cytoskeletal remodeling of the LC cells. Interestingly, blocking ILK in TM cells results in reduced cell spreading, actin polymerization, and the localization of talin and ILK in focal adhesions (FAs). This suggests that in POAG ONH astrocytes,

Figure 1. Signaling pathway activation profiles in glaucoma. Pathway activation strength (PAS) values were calculated by processing transcriptomic data obtained in human trabecular meshwork samples (datasets GSE4316 (A) and GSE27276 (B)) or lamina cribrosa samples (data sets GSE45570 (C) and GSE13534 (D)) using the AMD Medicine software suite. The fifty most dysregulated pathways compared to normal controls are shown. Blue bars represent PAS averages for each pathway denoting the degree of up regulation or down regulation. PAS presented on this figure passed the following filters PAS < 1.5 and PAS > 1.5 in all 4 datasets.
changes in ILK activity induced by the elevated IOP levels, may affect actin cytoskeleton organization and contractility in the TM.21

p38 is involved in the disruption of glucocorticoid receptor (GR) activity leading to a reduced GR nuclear translocation and signal transduction.22 Correspondingly, an up-regulation of the p38 pathway was seen 48 hours post-exposure, which paralleled a down-regulation in the GR pathway (activated by inflammatory cytokines) (Fig. 2). Since glucocorticoids mediate anti-inflammatory signaling pathways, their signaling may be down-regulated to facilitate the inflammatory responses generated from optic neuropathy associated with POAG.

The Wnt main pathway is up-regulated after 24 hours of exposure but then down-regulated after 48 hours along with the Wnt/PKC and Wnt/CREB3 pathways (Fig. 2). Dysregulation in the Wnt pathway is known to contribute to retinal degeneration, cataract, exudative vitreoretinopathy, ocular tumor, and several congenital eye disorders.23 Moreover, myocilin may act as a modulator of Wnt signaling through interactions with the Fzd family of Wnt receptors and antagonists of Wnt signaling like sFRPs and WIF-1.23 Therefore, in the case of myocilin linked-POAG, elevated myocilin levels may also contribute to dysregulation of the Wnt pathway. Overexpression of the Wnt antagonist sFRP1 in the mouse eye caused rapid elevation of IOP,23 suggesting that down-regulation of Wnt signaling may underlie the glaucomatous phenotype.

**Figure 2.** Effect of increased pressure on signaling pathway activation in cultured lamina cribrosa cells. Pathway activation strength (PAS) values were calculated by processing and analyzing the data set GSE758 using the AMD Medicine software suite to understand the signaling pathway profile associated with elevated pressure in lamina cribrosa. Blue bars, Orange bars and Gray bars represent PAS averages and the degree of pathway up-regulation or down-regulation after 6 hour, 24 hour and 48 hour exposure to elevated pressure, respectively.

**Signaling pathways activated in MYOC mutants**

Myocilin (MYOC), a protein with cytoskeletal function, is highly expressed in TM, ciliary body and can be found in AH.24,25 MYOC mutations are responsible for 3–4% of adult-onset POAG.24,26 Wild type MYOC is processed and secreted into TM ECM where it interacts with fibronectin whereas mutated MYOC can accumulate in cells and can impair the function of the TM, thereby resisting AH outflow and increasing IOP.24,25 More than 70 types of MYOC mutations are
associated with glaucoma each having a slightly different disease phenotype.26

In order to identify unique and common pathways between cells bearing wild type MYOC and mutant forms of this protein we processed data sets E-MEXP-3427, E-MEXP-3435, E-MEXP-3434 and E-MEXP-3439, representing transcriptional profiles of 4 TM cell lines each expressing 4 different MYOC mutations Q368X, R342K, D380N and K423E, and compared them to the control dataset E-MEXP-3440 of a TM cell line expressing wild type MYOC. Q368X is the most common mutant type (29% of the variants) with a mild phenotype. Although mutants R342K, D380N and K423E are much less abundant, they result in a severe clinical outcome. A Venn diagram (Fig. 3) demonstrates that only 2 pathways (upregulation of the caspase cascade pathway and down-regulation of STAT3 pathway related to growth arrest and differentiation) were commonly dysregulated in both mutant and wild-type MYOC-bearing cells.

TM cells containing high levels of MYOC are highly sensitive to apoptosis.27 As mentioned above, increased expression of MYOC results in loss of actin stress fibers and focal adhesions.27,28 MYOC may decrease cellular adhesion to fibronectin, compromising cellular homeostasis, thereby inducing susceptibility to apoptosis.28 This may explain the up-regulation on the caspase cascade revealed by our analysis.

Subsequently, a parallel decrease in STAT3 pathway related to growth and differentiation may be associated with the up-regulation of pro-apoptotic genes within the caspase cascades, leading to a pro-apoptotic fate of the cells.29 Notably, the p38 signaling pathway was up-regulated in all MYOC mutants cell lines compared to the wild type control (Fig. 3).

**TGFβ treated LC samples and glaucomatous LC samples cluster together at the pathway level**

In POAG, the ECM remodeling of the LC is believed to be regulated by high levels of TGFβ and eventually lead to a fibrotic ONH.30 To further delineate the intricate signaling pathways involved in the ECM remodeling of the LC, we have analyzed data sets GSE2378 and GSE2705 (containing glaucomatous astrocytes treated or untreated with TGFβ). To directly compare pathways activated in glaucomatous LC, TGFβ-treated LC and control LC, we have created the hierarchically clustered heat-map of differentially activated pathways in all 3 sample groups (Fig. 4). Interestingly, pathways dysregulated in TGFβ-treated cells, cluster together with pathways disturbed in glaucomatous LC, but not with the normal controls (Fig. 4). Notably, pro-survival pathways up-regulated in TGFβ-treated and glaucomatous LC (such as JAK-STAT, PAK, JNK and AKT), are highly associated with pro-fibrotic processes orchestrated by TGFβ.31,32 further supporting the idea that TGFβ induced fibrogenesis is an integral part of the POAG development.

**Discussion**

Although POAG is a leading cause of blindness worldwide, the molecular mechanisms underlying its initiation, maintenance and progression are not yet fully understood and remain to be elucidated. Here we applied AMD Medicine, a new bioinformatics software suite, for qualitative analysis of intracellular signaling pathway activation using transcriptomic data, to assess the network of molecular signaling associated with the POAG phenotype.

It is well documented that AH outflow blockage in glaucomatous eyes results from the resistance created by changes in the quality and amount of the ECM in the juxtacanalicular region of the TM33 (Fig. 5). These changes are believed to be highly associated with overexpression of TGFβ, which was reported to orchestrate the pro-fibrotic signaling that may affect the TM ECM turnover.33 TGFβ facilitates net matrix deposition by over-production of fibronectin (FN), collagen I and IV and inhibitors of ECM proteases, such as plasminogen activator inhibitor-1 (PAI-1) and tissue inhibitors of matrix metalloproteinases (TIMPs) that degrade matrix components.34,35 Subsequently, abrupt changes in ECM composition may lead to disrupted cell-cell signaling and non-specific interactions, resulting in apoptotic cell death.35

**Figure 3.** p38 signaling activation is associated with myocilin (MYOC) mutations induced glaucoma. Datasets E-MEXP-3427, E-MEXP-3435, E-MEXP-3434, E-MEXP-3439, containing gene expression profiles of 4 TM cell lines with different MYOC mutations and control data set (E-MEXP-3440) were processed and analyzed using AMD Medicine software suite. The non-intersecting blue region shows 1 pathway that was differentially activated in cell lines bearing 4 different MYOC mutants only and the non-intersecting green region shows 315 pathways that were activated in cell lines overexpressing wild type MYOC. The intersection of the blue and green circle represents pathways shared between WT MYOC and mutants with similar PAS values; the caspase cascade that is up-regulated and the STAT3 pathway that is down-regulated.
Our pathway activation analysis has demonstrated a significant AKT pathway up-regulation in POAG TM samples compared to normal controls. The AKT signaling main pathway is an important regulator of cell survival in response to growth factors and other extracellular stimuli and, when activated, exerts anti-apoptotic activity by preventing the release of cytochrome c from mitochondria. Since glaucomatous pathophysiology involves decrease in TM cellularity due to apoptosis,

Figure 4. TGFβ treated LC cells and glaucoma affected LC cells cluster together at the pathway level. Datasets GSE2378 and GSE3705 (containing glaucomatous astrocytes treated or untreated with TGFβ) were processed and analyzed using the AMD Medicine software suite. To directly compare pathways activated in glaucomatous LC, TGFβ-treated LC and control LC, we have created the hierarchically clustered heatmap of differentially activated pathways in all 3 sample cohorts. Red boxes represent pathway up-regulation and blue boxes represent pathway down-regulation. PAS values generated for in-vitro TGFβ treated human LC cells substantially correlate with PAS values obtained for glaucomatous LC cells and numerous pathways dysregulated in these 2 cohorts cluster together, but not with the normal controls, suggesting that TGFβ induced fibrogenesis is an integral part of glaucoma development.

Figure 5. Trabecular meshwork and lamina cribrosa are 2 main players in glaucoma progression. Model of eye tissues associated with glaucoma pathophysiology. (A) Diagram of eye tissues involved in glaucoma pathogenesis – 2 blue rectangles: eye angle containing trabecular meshwork (TM) and optic nerve head containing lamina cribrosa (LC) (B). Eye angle containing trabecular meshwork located between the cornea and the iris. Aqueous humor (AH) is produced by the ciliary body in the posterior chamber, flows into the anterior chamber and is finally drained into the Schlemm’s canal (SC) via TM. Arrows show direction of AH movement. Blue rectangle corresponds to the larger trabecular meshwork insert. (C) Trabecular meshwork insert showing AH flow (arrows) though juxtacanalicular tissue into Schlemm canal. AH outflow blockage due to clogging of TM results in elevated IOP. (D) Optic nerve head containing axons of RGC and lamina cribrosa structure. (E) Insert showing fine morphology of collagen fibers of lamina cribrosa. Lamina cribrosa ECM proper construction is vitally important for LC function. Misalignment of collagen fibers due to ECM rearrangement results in loss of mechanical resistance of LC. (F) SEM of trabecular meshwork (re-print from 73 with publisher permission). (G) SEM of lamina cribrosa (re-print from 74 with publisher permission).
RGCl death from the optic neuropathy and ECM remodeling. AKT activation might play a protective role against decrease in TM cellularity. Correspondingly, the substantial down-regulation of the PTEN main pathway (a negative AKT regulator) and up-regulation of the PAK main pathway (PAK acts as a scaffold to facilitate AKT stimulation by PDK1 and aids in recruitment of AKT to the membrane), seen in POAG samples, may further support these suggestions.

In addition to AKT, our analysis revealed up-regulation in other pro-survival signaling pathways in POAG TM, such as the p38 signaling main pathway, extracellular receptor kinases (ERK) and c-jun N-terminal kinases (JNK). p38 function is neuroprotective and its up-regulation facilitates RGC survival after ischemia or reperfusion injury. Moreover, it has been reported that ERK, p38 and JNK signal transduction pathways are relatively unresponsive in glaucomatous TM cells as compared to normal cells, suggesting their constitutive signaling activity in POAG. Additionally, through silencing of the pro-apoptotic gene BAD, and induction of the anti-apoptotic gene BCL-2, MAPK and AKT pathways (which were up-regulated in ONH) promote neuronal survival by rescuing RGC from death occurring from optic nerve injury. Interestingly, the p38 pathway related to cell motility, inflammation, apoptosis and osmoregulation was up-regulated in the cell lines over-expressing mutated MYOC. Because the MYOC expression pattern is complex, its biological function in normal and glaucomatous eyes has been difficult to elucidate. However, data indicate that POAG caused by mutations in MYOC may result primarily from protein misfolding or improper protein trafficking and its expression is also induced by TGFβ. Additionally, MYOC is associated with the downregulation of RhoA, which also can facilitate phosphorylation of p38 and activation of MAPK related pathways.

The ILK signaling main pathway, which facilitates the increase in matrix metalloproteinases (MMP-2), was also up-regulated in POAG samples compared to normal controls. Moreover, ILK pathway related to migration and vasculogenesis was significantly up-regulated in ONH astrocytes after 48 hr exposure to HP. Interestingly, in glaucomatous conditions, TM cells sense mechanical stretching and respond by increasing levels of MMP and tissue inhibitor of matrix metalloproteinase (TIMP) through integrin–ECM interactions to reverse outflow resistance and re-establish IOP homeostasis. Therefore, it is possible that ILK protects against mechanical stretching of the ocular tissue from AH outflow blockage. Notably, several studies have shown that ILK is also involved in the activation of ERK and p38 MAPK signaling during the development of hepatic and pulmonary fibrosis, suggesting that similar signal transduction cascades may also occur in POAG.

The MAPK/ERK signaling cascade is a major pathway controlling cellular processes associated with fibrogenesis, including growth, proliferation, and survival whereas PAK/P38 signaling plays a key role in pro-fibrogenic epithelial–mesenchymal transition (EMT) and was shown to be up-regulated during organ fibrogenesis. Moreover, ERK, p38 MAPK and ILK pathways are principally related to TGFβ signaling, which plays a central role in fibrotic disorders by inducing multiple pro-fibrogenic and immunosuppressive effects in various distinct organs including the eye. In glaucoma, TGFβ-induced fibrotic processes are known to orchestrate the build-up of ECM materials in the TM at the anterior of the eye, and in the LC at the ONH. For example, GFAP-negative LC cells constitutively release TGFβ-induced pro-fibrotic factors, such as connective tissue growth factor (CTGF) and platelet-derived growth factor-alpha (PDGF-alpha). Moreover, TGFβ induces expression and release of collagen type IA1, α-smooth muscle actin (αSMA), and alters cell–ECM interaction rigidity in POAG LC cells. Notably, we found that various pro-survival pathways highly associated with pro-fibrotic processes such as the AKT Signaling Main Pathway, CXC Chemokine Receptor Pathway, EGF Main Pathway, ERK Signaling Main Pathway, Hedgehog Signaling in Mammals Main Pathway, IGF1R Signaling Main Pathway, IP3 Main Pathway, JAK–STAT Main Pathway and JNK Main Pathway, were up-regulated in TGFβ-treated glaucomatous astrocytes and glaucomatous LC samples. Although these data support previous observations that TGFβ-induced fibrogenesis is an integral part of the POAG, these TGFβ-induced pro-survival pathways may have a neuroprotective role against ECM remodeling and/or IOP stress-related apoptosis. These pathways may also facilitate progenitor cell proliferation and astrocyte differentiation to cope up with the damage from fibrosis.

In contrast, the cognac synthase kinase 3 (GSK3) pathway was down-regulated in POAG TM. Down-regulation of GSK3 has been shown to induce expression of active β-catenin and to suppress levels of certain ECM proteins, even under stimulatory effects of TGFβ. Consequently, down-regulation of GSK might play a role in activation of canonical WNT signaling by statins (HMG-CoA reductase inhibitors) and facilitate AH outflow to re-establish IOP homeostasis under glaucomatous conditions. Interestingly, emerging evidence has shown that the cross-talk between WNT and TGFβ signaling plays important roles in TM
homeostasis and IOP regulation.\textsuperscript{66} While only a few genes that mediate the crosstalk between the 2 pathways are currently discovered, manipulating these mediators may provide a more effective way of restoring aqueous outflow in the TM, and possibly treating glaucoma.

**Conclusion**

Malfunction of 2 fine structures in the human eye (TM and LC) could lead to glaucoma since the ability to carry their function heavily depends on morphology,\textsuperscript{67} which can be easily affected by pro-fibrotic ECM rearrangements (Fig. 5). As the main function of TM is to filter out AH, slight changes in ECM can cause clogging of the fine TM filtering system (Fig. 5B, C, G). The main function of LC is to withstand IOP provided by precise configuration of collagen fibers, which when misaligned can cause loss of mechanical resistance and subsequent atrophy of ONH (Fig. 5D, E, F). Only one third of people with glaucoma have normal or near normal IOP, whereas one third of patients with high IOP do not develop glaucoma,\textsuperscript{68,69} which represents the main challenge in understanding glaucoma biology. While elevated IOP is a major risk factor for glaucoma, ECM remodeling of the LC as a result of pro-fibrotic processes can provide a mechanistic explanation for the glaucoma phenotype in a considerable portion of patients with normal IOP. Simultaneously, LC resistance to ECM rearrangement affecting LC ability to withstand elevated IOP could explain why some people with increased ocular pressure do not develop the glaucoma phenotype (Fig. 6).

Although our results are correlations, and future confirmatory studies are warranted to validate these observations, it is tempting to speculate that until increased IOP gets counter balanced by a fibrosis resistant LC (probably thicker and smaller in diameter that can withstand elevated IOP\textsuperscript{70}), patients will most probably remain POAG free. However, when the LC collagen structure is affected by fibrosis and LC mechanical resistance drops below a certain threshold, it may create a favorable condition for ONH cupping and deformation (glaucoma phenotype) even with healthy TB (normal IOP) (Fig. 6).

While it is well known that fibrosis plays a role in the glaucoma progression and a number of therapeutic approaches have been studied in an attempt to combat fibrosis, the fibrogenesis-driving mechanisms are not yet fully understood. Therefore, computational studies, like ours, may aid in uncovering complex molecular processes underlying the fibrotic changes in glaucoma which subsequently may lead to discovery of novel attractive therapeutic targets.

**Methods**

**Source datasets**

In this study, we utilized microarray gene expression data downloaded from NCBI GEO and ArrayExpress databases in order to examine pathways that are affected by glaucoma. The following data sets have been used: GSE4316, GSE27276, GSE45570, GSE13534, GSE758, E-MEXP-3440, GSE2378, GSE2705.

**Bioinformatics analysis and transcriptomic expression data pre-processing**

All microarray data pre-processing steps were performed in R version 3.1.0 using packages from Bioconductor. Microarray raw data were background adjusted and quantile normalized using the corresponding R packages. Obtained gene expression values were averaged across all replicates. Heat map generation and hierarchical clustering were performed using R package gplots. Statistical tests and correlation analysis were done with the MS Excel software.

**Signaling pathway analysis**

Preprocessed gene expression data were loaded into AMD Medicine pathology analysis software, a proprietary suite developed by Vision Genomic, Inc. which represents a cloud based implementation of the Oncofinder algorithm\textsuperscript{16,17,19,71} optimized for AMD/Glaucoma studies. As previously described,\textsuperscript{72} it enables calculation of the Pathway Activation Strength (PAS), a value which serves as a quantitative measure of differential pathway activation between the 2 states. PAS scoring is based on the expression level and the role of a particular gene in proprietary maps of 3 hundred of signaling pathways. Pathways with positive PAS values are considered up-regulated, while negative PAS values correspond to down-regulated pathways.

**Disclosure of potential conflicts of interest**

The authors declare no conflict of interest.

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