A Genome-Wide Scan for MicroRNA-Related Genetic Variants Associated With Primary Open-Angle Glaucoma

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PURPOSE: To identify microRNAs (miRNAs) involved in primary open-angle glaucoma (POAG), using genetic data. miRNAs are small noncoding RNAs that posttranscriptionally regulate gene expression. Genetic variants in miRNAs or miRNA-binding sites within gene 3′-untranslated regions (3′UTRs) are expected to affect miRNA function and contribute to disease risk.

METHODS: Data from the recent genome-wide association studies on intraocular pressure, vertical cup-to-disc ratio (VCDR), cup area and disc area were used to investigate the association of miRNAs with POAG endophenotypes. Putative targets of the associated miRNAs were studied according to their association with POAG and tested in cell line by transfection experiments for regulation by the miRNAs.

RESULTS: Of 411 miRNA variants, rs12803915:A/G in the terminal loop of pre-miR-612 and rs2273626:A/C in the seed sequence of miR-4707 were significantly associated with VCDR and cup area (P values < 1.2 × 10^-5). The first variant is demonstrated to increase the miR-612 expression. We showed that the second variant does not affect the miR-4707 biogenesis, but reduces the binding of miR-4707-3p to CARD10, a gene known to be involved in glaucoma. Moreover, of 72,052 miRNA-binding-site variants, 47 were significantly associated with four POAG endophenotypes (P value < 6.9 × 10^-6). Of these, we highlighted 10 variants that are more likely to affect miRNA-mediated gene regulation in POAG. These include rs3217992 and rs1063192, which have been shown experimentally to affect miR-138-3p– and miR-323b-5p–mediated regulation of CDKN2B.

CONCLUSIONS: We identified a number of miRNAs that are associated with POAG endophenotypes. The identified miRNAs and their target genes are candidates for future studies on miRNA-related therapies for POAG.

Keywords: primary open-angle glaucoma (POAG), microRNA, genetic variant, GWAS

Primary open-angle glaucoma (POAG), the most common optic neuropathy, is the leading cause of irreversible blindness, affecting approximately 60 million individuals worldwide.1,2 The disease is characterized by progressive loss of retinal ganglion cells and optic nerve degeneration that can be secondary to elevated intraocular pressure (IOP).3 The optic nerve damage is characterized by an increase in cup size, which is the central area of the optic disc. Cup enlargement can be measured by the vertical cup-to-disc ratio (VCDR), comparing the vertical diameter of the cup with vertical diameter of the total optic disc.4 The VCDR ranges from 0 to 1; a ratio above 0.7 or an asymmetry between eyes above 0.2 is considered as suspect for glaucoma in the clinical setting.5 POAG is presumed to be a complex progressive neurodegenerative disorder caused by multiple genetic as well as environmental factors.6 Previous genome-wide association studies (GWASs) have revealed a number of susceptibility loci for POAG by studying the disease directly or its endophenotypes including IOP and optic disc parameters (VCDR, cup area, and disc area).6,5 Most of the associated variants identified by GWASs are located in noncoding regions of the genome and their mechanistic contributions to POAG and its endophenotypes remain poorly understood.5,6

MicroRNAs (miRNAs) are small noncoding RNAs, consisting of 19 to 22 nucleotides, that posttranscriptionally regulate gene expression.7 There are strong indications that miRNAs play important roles in the pathogenesis of POAG.8-11 For example, miR-29b and miR-24 are involved in gene regulation in trabecular meshwork cells.12 Moreover, the miRNA expression levels have been linked to maintaining the balance of the aqueous humor, the change in the trabecular meshwork, and the apoptosis of the retinal ganglion cells.13-15 A number of miRNAs (e.g., miR-200c, miR-204, miR-183, and miR-182) are also reported as potential diagnostic biomarkers or therapeutic
targets for glaucoma.\textsuperscript{9,16} The biogenesis of miRNAs is a multistep coordinated process.\textsuperscript{17,18} In the nucleus, miRNA genes are initially transcribed as long primary transcripts. Further processing and cleavage by the RNase \textit{Drosha} and \textit{Dicer} enzymes generate mature miRNAs.\textsuperscript{17,18} The mature miRNA is subsequently incorporated into the RNA-induced silencing complex (RISC) to interact with the 3\textsuperscript{\prime}-untranslated region (3\textsuperscript{\prime}UTR) of target mRNAs, resulting in mRNA degradation or translational repression.\textsuperscript{7,18} Genetic variants in miRNA-encoding sequences can have profound effects on miRNA biogenesis and function.\textsuperscript{19,20} In addition, polymorphisms located in miRNA-binding sites within the 3\textsuperscript{\prime}UTR of target genes are expected to affect miRNA-mediated gene regulation.\textsuperscript{20,21} Previous studies\textsuperscript{19,21–25} have shown that such miRNA-related variants contribute to complex disease risk; a candidate variant association study\textsuperscript{24} has also recently reported the association between genetic variation in miR-182 and POAG. In this study, we applied an in silico study on the existing GWAS data of glaucoma endophenotypes and were included for association analysis. The Bonferroni correction was used to adjust the \textit{P} value for multiple comparisons. To this end, we divided the critical \textit{P} value (\(\alpha = 0.05\)) by the number of genetic variants in each group (miRNAs and miRNA-binding sites). As we studied 411 variants in miRNAs, the significance threshold was set on \(1.22 \times 10^{-4} (0.05/411)\) for miRNA genetic variants.

Likewise, the significance threshold for genetic variants in miRNA-binding sites was set on \(6.94 \times 10^{-7} (0.05/72,052)\), as we studied 72,052 variants in this group (Fig. 1). Regional association plots showing the association of miRNA-related variants with the studied phenotypes were made using LocusZoom.\textsuperscript{27}

### Methods

#### Genome-Wide Association Studies on Glaucoma Endophenotypes

To examine the association of miRNA-related genetic variants with POAG endophenotypes, we used data from the recent GWAS on glaucoma endophenotypes provided by the International Glaucoma Genetics Consortium (IGGC).\textsuperscript{5} Characteristics of the IGGC have been described elsewhere.\textsuperscript{5} In brief, the IGGC has conducted genome-wide association meta-analyses on four glaucoma endophenotypes including IOP (\(n = 37,930\) individuals), VCDR (\(n = 23,899\) individuals), cup area (\(n = 22,489\) individuals), and disc area (\(n = 22,504\) individuals). The IGGC, using imputation to the 1000 Genomes (1000G) reference panel, has tested the association of approximately 401,000 genetic variants in the GWAS data of glaucoma endophenotypes and were included for association analysis. These variants include 87 in pre-miRNA loci, 88 in mature miRNA sequences (19–22 nt), and miRNA seed regions (nucleotides 2–8 from the 5\textsuperscript{\prime}-end of a mature miRNA, which is defined as the most crucial part of an miRNA for target recognition and has to be perfectly complementary to the binding site of a target mRNA). Of 2420 miRNA variants, 411 (in 332 miRNAs) were selected for association analysis. These variants include 283 in pre-miRNA loci, 87 in mature miRNA sequences, and 41 in miRNA seed regions. Moreover, we extracted almost 401,000 genetic variants (including SNPs and INDELs) in miRNA-binding sites within the gene 3\textsuperscript{\prime}UTRs by using PolymiRTS v3 (database update October 2014).\textsuperscript{26} Of these, 72,052 miRNA-binding-site variants were available in the GWAS data of glaucoma endophenotypes and were included for association analysis. The Bonferroni correction was used to adjust the \textit{P} value for multiple comparisons. To this end, we divided the critical \textit{P} value (\(\alpha = 0.05\)) by the number of genetic variants in each group (miRNAs and miRNA-binding sites). As we studied 411 variants in miRNAs, the significance threshold was set on \(1.22 \times 10^{-4} (0.05/411)\) for miRNA genetic variants.

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#### Genetic Variants Located in miRNAs and miRNA-Binding Sites

A total of 2420 genetic variants located in miRNA-encoding sequences were retrieved from miRNASNP v2 (database update April 2015)\textsuperscript{28} and literature review (searching for miRNA genetic variants in PubMed). These include variants in precursor miRNAs (60–80 nucleotides [nt]), mature miRNA sequences (19–22 nt), and miRNA seed regions (nucleotides 2–8 from the 5\textsuperscript{\prime}-end of a mature miRNA, which is defined as the most crucial part of an miRNA for target recognition and has to be perfectly complementary to the binding site of a target mRNA). Of 2420 miRNA variants, 411 (in 332 miRNAs) were available in the GWAS data of glaucoma endophenotypes and were selected for association analysis. These variants include 283 in pre-miRNA loci, 87 in mature miRNA sequences, and 41 in miRNA seed regions. Moreover, we extracted almost 401,000 genetic variants (including SNPs and INDELs) in miRNA-binding sites within the gene 3\textsuperscript{\prime}UTRs by using PolymiRTS v3 (database update October 2014).\textsuperscript{26} Of these, 72,052 miRNA-binding-site variants were available in the GWAS data of glaucoma endophenotypes and were included for association analysis. The Bonferroni correction was used to adjust the \textit{P} value for multiple comparisons. To this end, we divided the critical \textit{P} value (\(\alpha = 0.05\)) by the number of genetic variants in each group (miRNAs and miRNA-binding sites). As we studied 411 variants in miRNAs, the significance threshold was set on \(1.22 \times 10^{-4} (0.05/411)\) for miRNA genetic variants.

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#### The Effect of Variants on the miRNA Secondary Structure and Expression

The secondary structure of pre-miRNA is critical for the miRNA biogenesis.\textsuperscript{28} The Vienna RNAfold algorithm (ViennaRNA package 2.0) was used to predict the impact of variants in miRNAs on the hairpin stem-loop structure of pre-miRNAs.\textsuperscript{29} This program calculates the changes in minimum free energy (MFE) of the thermodynamic ensemble of the hairpin structure of miRNA (wild type and mutant). Next, to test the functional impact of an miRNA variant on the expression of the mature miRNA, we cloned the pre-miRNA sequence containing either the major or minor allele behind the gene encoding green fluorescent protein (GFP) in the expression plasmid MSCV-BC (Murine Stem Cell Virus-Bar Coded), resulting in GFP-miRNA fusion transcripts as described previously.\textsuperscript{30} For cloning purposes, the restriction enzyme sites \textit{XhoI} and \textit{EcoRI} were inserted in forward and reverse primers, respectively. The primers are listed in Supplementary Table S1. We used Sanger sequencing to validate the inserts of all constructs. HEK293 cells were plated at a density of \(5 \times 10^{4}\) cells per well in a six-well plate and were cultured in 2 mL Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal bovine serum and 100 units/mL penicillin/streptomycin in a 37\textdegreeC incubator with 5% \textit{CO}_2. The next day, cells were transiently transfected with 1 \(\mu\)g GFP-miRNA constructs containing either the major or minor allele by using Lipofectamine\textsuperscript{R} RNAiMAX (Invitrogen, Carlsbad, CA, USA) according to the manufacturer's instructions. After 48 hours, total RNA was isolated with the TRIzol reagent (Invitrogen) according to the manufacturer’s protocols. The concentration and purity of RNA samples were determined with a NanoDrop ND-1000 spectrophotometer (NanoDrop, Wilmington, DE, USA). TaqMan Assays (Applied Biosystems, Foster City, CA, USA) were used to measure the expression levels of GFP and miRNA by quantitative RT-PCR. The expression levels of mutant and wild-type miRNA relative to GFP were calculated as previously described.\textsuperscript{30} RNU6B was used as an internal loading control for miRNA expression. All experiments were performed in triplicate and repeated at least three times.

#### Association of miRNA Target Genes With Glaucoma Endophenotypes

The biological role of miRNAs is dictated through regulating expression of their target mRNAs. We aimed to identify target genes that may mediate the downstream effects of the associated miRNAs in relation to POAG endophenotypes. Since such genes should be also associated with the studied phenotypes, we used the GWAS data in a candidate gene approach and searched among the putative target genes of an miRNA for those that were associated with POAG endophenotypes. To this end, we extracted the miRNA putative targets by using two commonly used online databases: TargetScan v7.0 (total context \textit{\&} score > 0.13)\textsuperscript{31} and miRDB.\textsuperscript{32} TargetScan (http://www.TargetScan.org/) (in the public domain) predicts
biological targets of miRNAs by searching for the presence of conserved sites that match the seed region of each miRNA. The predictions in this database are ranked by the predicted efficacy of targeting as calculated by using cumulative context scores of the sites by considering several features to predict the most effective target mRNAs.\textsuperscript{31} The targets in miRDB (http://mirdb.org//; in the public domain) were predicted by a bioinformatics tool, which was developed by analyzing thousands of miRNA–target interactions from high-throughput sequencing experiments.\textsuperscript{32} The GWAS data from glaucoma endophenotypes were used to examine the association of genetic variants in the predicted miRNA target genes with the phenotype of interest.\textsuperscript{5} The Bonferroni correction was used to calculate the significance level based on the number of tested genetic variants in all predicted target genes of each miRNA.

Interaction Analysis Between an miRNA and Its Target Genes Using the Rotterdam Study Data

The external validity of association between an miRNA variant and associated phenotype may improve when the variant affects the miRNA target genes involved in the phenotype. We thus tested the interaction between associated miRNA variants and the most significant variant in their related target genes in relation to POAG endophenotype, using the Rotterdam Study data. To examine the interaction, we introduced the statistical product of miRNA variant and target gene variant in the logistic regression model, adjusted for age and sex: POAG endophenotype \( \sim \) age + sex + miRNA SNP + top SNP in target gene + (miRNA SNP \times\) top SNP in target gene). The design of the Rotterdam Study has been described in detail elsewhere.\textsuperscript{33} The baseline characteristics of the subjects are shown in Supplementary Table S2.

Luciferase Reporter Assays

A luciferase reporter assay system was used to examine the interaction between an miRNA and the 3’UTR of its putative target gene. In addition, this experiment was used to determine the impact of candidate variant on the miRNA-target gene interaction. To this end, primers were designed to amplify the 3’UTR sequence of the miRNA target gene and included restriction enzyme sites XbaI for the forward primer and ApatI for the reverse. The 3’UTR sequences containing putative binding site (wild type and mutated) of the miRNA were amplified and cloned into the pGL3 Luciferase reporter vector (Promega, Madison, WI, USA) downstream of the Luciferase open reading frame.\textsuperscript{30} All the primers are shown in Supplementary Table S3. The inserts of all constructs were...
confirmed by Sanger sequencing. HEK293 cells (n = 10,000) were plated into 96-well plates and cotransfected with 1 μg pG5s containing the 3′UTR with either the major or minor allele, miRNA mimic (mirVana Mimics; Thermo Fischer Scientific, Waltham, MA, USA), and a plasmid expressing the Renilla luciferase that served as transfection control, with Lipofectamine RNAiMAX (Invitrogen). Luciferase activity was measured with the Dual-Glo Luciferase Assay System according to manufacturer’s protocol (Promega). Renilla activity was used for normalization of the data. All experiments were performed in triplicate and repeated in three independent experiments.

Functional Annotation of miRNA-Binding-Site Variants Associated With Glaucoma Endophenotypes

Specific criteria have been suggested to establish whether 3′UTR variants located in miRNA-binding sites may affect miRNA-mediated gene regulation.20,21 These include association between the variant and the phenotype of interest, expression of host gene and related miRNA in a relevant tissue, and an allele-specific regulation of the target transcript by miRNA.20,21 These criteria were used to prioritize the miRNA-binding-site variants that are more likely to be functional. We first retrieved proxy SNPs in high linkage disequilibrium (LD) (R² threshold > 0.8, limit distance 100 kb, and population panel CEU) with the binding-site variants and checked their effects on protein structure, gene regulation, and splicing by using the HaploReg v4.1 (http://www.broadinstitute.org/mammals/haploreg/haploreg.php; in the public domain). HaploReg 4.1 was also used to evaluate the functional potential of the LD variants associated with glaucoma endophenotypes. To scan the correlation between the identified variants and expression levels of the host transcripts, we used expression quantitative trait loci (cis-eQTL) data from the GTExV6 (http://www.gtexportal.org/home/; in the public domain) and Genenetwork (http://genenetwork.nl/bloodedqtlbrowser/; in the public domain). To check whether miRNAs in our collection are expressed in ocular tissues, two online databases were screened, miRBase54 and HMDD.55 Moreover, we used expression data from previous miRNA profiling studies in human ocular tissues, including ciliary body, cornea, and trabecular meshwork and aqueous humor.13-15 Other miRNA information, including miRNA conservation in different species, was obtained from miRBase (release 20).20 In addition, the Ocular Tissue Database (https://genome.uiowa.edu/otdb/; in the public domain) was used to examine the expression of the identified miRNA target genes across the eye tissues. Supplementary Table S4 shows a list of web tools and databases that we used for our analyses.

RESULTS

Two miRNA Variants Were Associated With POAG Endophenotypes

We examined the association of 411 miRNA variants (in 332 miRNA genes) with IOP VCDR, cup area, and disc area (Fig. 1). Two miRNA variants passed the Bonferroni-corrected significance threshold of 1.22 × 10⁻⁴ (0.05/411). These variants include rs12803915:A/G (Chr11:65444508) in the terminal loop of pre-miR-612 sequence associated with VCDR (P value = 4.6 × 10⁻⁴, β = 0.009) and cup area (P value = 1.2 × 10⁻⁵, β = 0.004), and rs2273626:A/C (Chr14:22956973) in the seed region of pre–miR-4707 sequence associated with VCDR (P value = 5.8 × 10⁻⁵, β = 0.005) and cup area (P value = 9.9 × 10⁻⁵, β = 0.008) (Table 1). No miRNA variants were significantly associated with IOP and disc area. Supplementary Table 55 shows genetic variants in miRNAs that are nominally associated (P value < 0.05) with POAG endophenotypes.

The Impact of Variants on the miRNA Secondary Structure and Expression

The impact of rs12803915 and rs2273626 on the miRNA biogenesis was measured by performing structural analysis of the miRNA hairpin, using Vienna RNAfold algorithm.56 We observed 0.8 kcal/mol difference in MFE of the thermodynamic ensemble of the miRNA hairpin structure for pre–miR-612, suggesting that the variants may affect their miRNA production (Supplementary Fig. S1). Rs12803915 (minor allele A) in pre–miR-612 has been demonstrated previously to increase expression of the mature miRNA (Fig. S1). Rs2273626 (minor allele T behind GFP) in the pMSCV-BC vector. HEK293 cells transfected with the pre-miRNA sequence containing the rs2273626 major allele G or the minor allele T behind GFP in the pMSCV-BC vector. HEK293 cells transfected with the pre-miRNA. We observed no significant difference between expression levels of mature miR-4707 in cells transfected with the pre-miRNA sequence containing the major and the minor allele (Fig. 2A). This indicates that the effect of rs2273626 on the miR-4707 biogenesis is minor and not detectable with this experimental setup.

### Table 1. Association of Top miRNA Variants With POAG Endophenotypes

| Phenotype | SNP ID     | Chr | Position | A1/A2 | MAF   | miRNA ID | SNP Position | MFE   | P Value  | β     | SE    |
|-----------|------------|-----|----------|-------|-------|-----------|--------------|-------|----------|-------|-------|
| VCDR      | rs12803915 | 11  | 65444508 | A/G   | 0.14  | miR-612  | Stem region  | 0.8   | 4.6 × 10⁻⁴ | −0.009 | 0.0015|
| IOP       | rs2273626  | 14  | 22956973 | A/C   | 0.48  | miR-4707 | Seed         | 5.8   | 9.5 × 10⁻⁵ | −0.005 | 0.0012|

*Shown are top three miRNA variants with the most significant association with each of the POAG endophenotypes. Two variants that pass the significant threshold (P value < 1.22 × 10⁻⁴) are shown in bold. A, allele; Chr, chromosome; SE, standard error; MFE, change in minimum free energy.*
Target Genes of miR-612 and miR-4707 Associated With VCDR and Cup Area

To identify target genes that may mediate the downstream effect of miR-612 and miR-4707 in relation to VCDR and cup area, we studied the association of their putative target genes with these phenotypes, using the GWAS data. Supplementary Table S6 displays four predicted target genes of miR-612 and eight predicted target genes of miR-4707 that are associated with VCDR and cup area. Using data from the Ocular Tissue Database, we showed that these associated target genes are expressed in the eye tissues (Supplementary Table S7). We further tested the interaction between the identified miRNA variants and the top variant in their highlighted predicted target genes (associated with glaucoma endophenotypes and expressed in the eye), using the Rotterdam Study data. This analysis showed a significant interaction ($P = 0.028$) between the relative luciferase activity of the CARD10 3' UTR construct in the presence of miR-4707-3p containing the major allele and the minor allele. Our results suggest that rs2273626 diminishes the regulatory interaction between miR-4707-3p and CARD10, resulting in increased CARD10 levels. All experiments were performed in triplicates and repeated at least three times. Error bars represent standard deviation (SD). NS, nonsignificant.

Multiple miRNA-Binding-Site SNPs Were Associated With POAG Endophenotypes

In the second part of our study, we examined the associations of 72,052 miRNA-binding-site SNPs (available in the GWAS data) with POAG endophenotypes (Supplementary Fig. S2). Of these, 47 SNPs (located in the 3' UTR of 21 genes) passed the Bonferroni-corrected significance threshold of $6.94 \times 10^{-7}$, which are associated with one or more of the POAG endophenotypes. These SNPs are predicted to affect the putative interaction between their host genes and a number of miRNAs that are shown in Supplementary Table S9. To highlight the binding-site variants that are more likely to affect miRNA-mediated gene regulation, we prioritized them on the basis of various criteria (e.g., the strength of association, LD pattern, eQTL data, coexpression of miRNA and target gene in relevant tissue). Haploreg v4.1 data, we found showed that many of the binding-site SNPs ($n = 37$) have no nonsynon-
**MicroRNA-Related Variants Associated With Glaucoma**

**UTR Variants Associated With POAG Endophenotypes Within miRNA-Binding Sites**

| rs1063192 | rs1132554 | rs10216533 | rs1057001 | rs6660601 | rs7344 | rs1052990 | rs1050285 | rs1043207 | rs8176751 |
|---------|---------|------------|-----------|-----------|--------|-----------|-----------|-----------|-----------|
| GWAS result | CA | VCDR | IOP | CA | IOP | IOP | VCDR | IOP | CA | 
| /C0 | 33 | 5.5 | 7 | 2.8 | 8 | 2.3 | 7 | 4.2 | /C0 | 11 | 4.0 | /C0 | 7 | 1.9 | /C0 | 9 | 2.9 | /C0 | 7 | 6.5 | /C0 | 7 | 1.1 |

**GWAS result**

Effect allele G, C. 

**Expression**

Gene ID: Table 2, Supplementary Fig. S3. These include two SNPs in the 3'UTR of CDKN2B, rs3217992 and rs1063192, which have been demonstrated previously to affect miR-138-3p- and miR-323b-5p-mediated regulation of CDKN2B.21

**DISCUSSION**

In this study, we performed a genome-wide scan to identify miRNAs associated with POAG endophenotypes, using genetic data. We found genetic variants in the miR-612 precursor and in the miR-4707 seed region significantly associated with VCDR and cup area. The variant in miR-612 has been previously demonstrated to increase the miRNA expression.27 We showed that the variant in miR-4707 does not influence the miRNA expression, but affects the binding of miR-4707 to one of its glucoma-associated target genes, CARD10. Moreover, we identified 47 SNPs in miRNA-binding sites (within the 3'UTR of 21 genes) that are significantly associated with POAG endophenotypes. After prioritization, we highlighted the 3'UTR variants that are more likely to affect miRNA-mediated gene regulation in POAG.

Several studies8,9,15 have shown previously the contribution of miRNAs to the pathophysiology of POAG. However, these studies are mainly focused on differentially expressed miRNAs identified by miRNA profiling in a small number of samples, which makes it difficult to interpret their results to the general population. In addition, miRNA profiling studies are sometimes subject to confounding bias or reverse causation, and cannot provide evidence for a causal role of the identified miRNAs in relation to a disease. Here, we used a genetic approach that is proven to be efficient and successful for identification of miRNAs involved in complex traits.19–23 In this approach, the starting point is a linkage between miRNA variants and the trait of interest. The main advantage of the genetic approach compared to miRNA expression profiling is that when a variant is associated with disease risk, it supports the idea that the miRNA may have a primary effect in the disease pathogenesis. Given that genetic variants are randomly inherited, it is unlikely that other extrinsic factors (e.g., lifestyle) are associated with them. Moreover, as the DNA sequence is constant over the life course, reverse causation is also refuted. Another advantage of the genetic approach is that diseases with no accessible affected tissues (e.g., eye tissue of glaucoma patients) can still be studied, since genomic DNA is available from other accessible tissues such as blood. The main limitation of the genetic approach is that it is restricted to miRNAs with genetic variants available in the GWAS data. Therefore, both genetic and miRNA profiling approaches are valuable and can work complementarily to analyze different aspects of the role of miRNAs in disease pathogenesis. The genetic approach indicates whether miRNAs are potentially causal or risk factors for a disease and the miRNA profiling approach demonstrates whether the expression of the disease-associated miRNAs is altered in the affected tissue.

We found two variants in the precursor and seed region of miRNAs significantly associated with POAG endophenotypes.
The first variant is rs12803915 located in miR-612. The GWAS data show that the variant minor allele A is associated with decreased VCDR and cup area. The variant minor allele has been demonstrated previously to increase the expression of mature miR-612 in vitro.\(^5\) The location of variant in the terminal loop of miR-612 possibly improves the pre-miRNA processing, and consequently increases the miRNA expression.\(^59-42\) Next, we highlighted four putative miR-612 targets (USH2A, FAM101A, JRK, and SIX4) that are also associated with VCDR and cup area and are expressed in the eye. These target genes might mediate the downstream effect of miR-612 in POAG. Together, our results suggest a protective role for miR-612 in relation to POAG. Future experimental studies are needed to further elucidate the role of miR-612 and its highlighted target genes in the pathogenesis of POAG.

The second associated variant is rs2273626 located in the seed sequence of miR-4707, an miRNA that is shown to be expressed in aqueous humor of glaucoma patients.\(^43\) The variant minor allele A was negatively associated with VCDR and cup area in the GWAS data.\(^4\) A polymorphism in an miRNA seed sequence is expected to strongly influence the miRNA activity and possibly risk of disease.\(^40,43\) We tested the two mechanisms through which a seed sequence variant may affect miR-4707 function. Our results showed that rs2273626 has no significant effect on the miRNA expression levels. However, the variant reduces the interaction between miR-4707-3p and CARD10, presumably resulting in higher CARD10 levels. The CARD10 gene is reported in several GWAS to be associated with glaucoma.\(^45-47\) The activity of CARD10 has been suggested to have a neuroprotective role in glaucomatous optic neuropathy.\(^45\) POAG is a disease of enhanced retinal ganglion cell apoptosis.\(^49\) It has been postulated that CARD10 controls optic disc area possibly through ganglion axonal survival via the activation of the NFXB signaling pathway.\(^49,50\) Overexpression of CARD10 has been shown to be related with increased cell survival and proliferation.\(^50,51\) Conversely, homozygous knockout of CARD10 in a mouse model can cause nonviability through neural tube defects.\(^52\) CARD10 has been also shown to be required in neural crest cell survival through G protein-coupled receptor induction of NFkB activation.\(^21,54\) Retinal ganglion cell damage from NFXB is likely to be due to glial cell activation and IL-1β secretion.\(^55\) Given the intimate and complex relationship between CARD10, NFXB, and apoptosis,\(^57\) it is biologically plausible that lower levels of CARD10 result in increased retinal ganglion cell apoptosis. In our study, the negative association between the miR-4707 variant and POAG may be explained in part by higher CARD10 expression due to the disruption of miR-4707-mediated gene regulation.

In the second part of this study, we investigated the association of miRNA-binding-site variants with POAG endophenotypes and identified 47 associated SNPs. GWASs are based on predefined criteria (e.g., the strength of association, LD tagging variants that are in high LD with functional variant(s) in the region; it is thus challenging to determine the exact localization of the variants that cause the associations. Here we prioritized the 47 associated SNPs on the basis of a set of predefined criteria (e.g., the strength of association, LD evaluation, eQTL analysis, coexpression of the miRNA and the target gene in relevant tissue).\(^20,21\) We highlighted 10 of the variants that are more likely to affect the putative miRNA-mediated regulation. Among them, rs1063192 and rs3217992 have been shown previously to affect the miRNA–mRNA interaction in vitro.\(^21,54\) These two SNPs reside in the 3'UTR of CDKN2B and are not in strong LD (R² > 0.8) with any other known CDKN2B variants. The CDKN2B gene encodes a cyclin-dependent kinase inhibitor and with its antisense (CDKN2B-AS1) lies in a well-known glaucoma-associated locus on Chr.19p21.\(^5,56\) The minor allele (G) at rs1063192 site is predicted to create a binding site for miR-323b-5p and the rs3217992 minor allele (T) is expected to disrupt the existing binding site of miR-138-2-3p within the 3'UTR of CDKN2B. Horwitz et al.\(^54\) have shown that these miRNAs control both CDKN2B mRNA and protein levels. Moreover, they have shown the correlation of rs1063192 minor allele with lower CDKN2B mRNA levels (improving the miR-323b-5p-mediated regulation) and conversely, the correlation of rs3217992 minor allele with higher CDKN2B mRNA levels (disrupting the miR-138-2p-mediated regulation) in adipose tissue.\(^54\) Although the impact of these variants on the CDKN2B expression needs to be demonstrated in the ocular tissues, the functional consequences of both variants on the miRNA-binding sites have been confirmed by Luciferase reporter assays in cell lines.\(^21,54\) Thus, an allelic-specific regulation of CDKN2B by miR-232b-5p and miR-138-2-3p may be considered as a potential mechanism, at least in part, to explain the association between CDKN2B (rs1063192 and rs3217992) and POAG.

In addition to the two experimentally validated variants in the 3'UTR of CDKN2B, we found several other 3'UTR variants associated with POAG endophenotypes that are located in miRNA-binding sites and are one of the top variants in their loci with no nonsynonymous proxy in high LD (R² > 0.8). These variants have the potential to be functional variants in their loci by affecting miRNA-mediated gene regulations and warrant further investigations. For example, the minor allele of rs1052990 in the 3'UTR of CAV2, a known gene for glaucoma, is predicted to disrupt the predicted binding site of miR-224-5p and increase the transcript levels of CAV2. The association of the rs1052990 minor allele with increased expression levels of CAV2 has been reported previously.\(^55\) In addition, both miR-224-5p and CAV2 are expressed in the eye.\(^56,57\) These data may suggest an allele-specific regulation of CAV2 by miR-224-5p as a functional mechanism underlying the observed GWAS association.

Although we believe our results are reliable, there are certain limitations that need to be considered in interpreting of our results. First, the reported miRNA-target interactions are predicted from in silico approach and need to be validated by experimental studies. Second, we used eQTL data from various tissues, but not eye tissue, to check the association between the identified 3'UTR variants and the expression of hosting genes. As gene expression and eQTL are tissue specific, in an optimal setting one should examine the associations in the ocular tissues. Finally, we were not able to perform the functional experiments in a relevant cell line for glaucoma disease, owing to slow growth rate and low transfection efficiency of retinal ganglion cells in culture. However, to gain insights into the impact of the identified variant on miR-4707, we performed our experiments with HEK293 cells, a commonly used cell line in this type of studies.

**CONCLUSIONS**

We systematically investigated the association of miRNA-related genetic variants with four POAG endophenotypes. We found variants in the miR-612 precursor and in the miR-4707 seed region that are significantly associated with VCDR and cup area. Our results showed that the minor allele of these variants may alter expression or targeting of the miRNAs. Further, we highlighted a number of 3'UTR variants associated with POAG endophenotypes that are located in the predicted miRNA-binding sites and may affect miRNA-mediated gene regulation in POAG. The identified miRNAs and target genes are candidates for future studies to determine their roles in the pathophysiology of POAG and their therapeutic potentials for glaucoma.
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**APPENDIX**

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