Metabolome-Wide Association Study of Primary Open Angle Glaucoma

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PURPOSE. To determine if primary open-angle glaucoma (POAG) patients can be differentiated from controls based on metabolic characteristics.

METHODS. We used ultra-high resolution mass spectrometry with C18 liquid chromatography for metabolomic analysis on frozen plasma samples from 72 POAG patients and 72 controls. Metabolome-wide Spearman correlation was performed to select differentially expressed metabolites (DEM) correlated with POAG. We corrected P values for multiple testing using Benjamini and Hochberg false discovery rate (FDR). Hierarchical cluster analysis (HCA) was used to depict the relationship between participants and DEM. Differentially expressed metabolites were matched to the METLIN metabolomics database; both DEM and metabolites significantly correlating with DEM were analyzed using MetaboAnalyst to identify metabolic pathways altered in POAG.

RESULTS. Of the 2440 m/z (mass/charge) features recovered after filtering, 41 differed between POAG cases and controls at FDR = 0.05. Hierarchical cluster analysis revealed these DEM to associate into eight clusters; three of these clusters contained the majority of the DEM and included palmitoylcarnitine, hydroxyergocalciferol, and high-resolution METLIN matches to sphingolipids, other vitamin D-related metabolites, and terpenes. MetaboAnalyst also indicated likely alteration in steroid biosynthesis pathways.

CONCLUSIONS. Global ultrahigh resolution metabolomics emphasized the importance of altered lipid metabolism in POAG. The results suggest specific metabolic processes, such as those involving palmitoylcarnitine, sphingolipids, vitamin D-related compounds, and steroid precursors, may contribute to POAG status and merit more detailed study with targeted methods.

Keywords: glaucoma, metabolomics, plasma, sphingosine, vitamin D, palmitoylcarnitine

Glaucoma is an insidiously progressive optic neuropathy that affects nearly 90 million people worldwide, making it a leading cause of irreversible blindness.¹ The high variation of incidence among races² and the variation in clinical manifestations of glaucoma³ emphasize the need to understand how specific genetic and/or metabolic differences among individuals significantly influence disease status. The risk of glaucoma is increased by multiple genetic factors susceptible to the influence of numerous environmental exposures.⁴ Mutations in MYOC⁵ and OPTN⁶ have been associated with juvenile open angle glaucoma and normal tension glaucoma, respectively, and have also been found in a small percentage of primary open angle glaucoma (POAG) patients. Genome-wide association studies have found links between common genetic variants and POAG, such as CAV1 and CAV2,⁷ as well as TMCO1 and CDKN2BAS.⁸ However, in total, these genetic variants linked to glaucoma are thought to account for approximately 5% of all POAG and normal tension glaucoma cases.⁹ More recent studies have revealed three new loci, ABCA1, APOE, and GMDS, to be associated with POAG,¹⁰ but the extent to which these associations involve the general population has yet to be characterized.

The presence of glaucoma has also been associated with metabolic factors, including an increase in plasma markers of oxidative stress, such as malondialdehyde (MDA) and conjugated diene,¹¹ as well as reduced total antioxidant status (TAS).¹² However, solitary genetic polymorphisms and single protein or metabolic biomarkers have been limited in their predictive capacity. These biomarker studies hint that future predictive measures are possible, but greater discovery of biomarkers associated with glaucoma is necessary to advance knowledge of the disease mechanism at a personalized level.

Ultrahigh resolution metabolic profiling using liquid chromatography-mass spectrometry (LC-MS) provides a practical approach to personalized medicine and has great potential to advance the evaluation of glaucoma risk. The ultrahigh mass resolution and mass accuracy of current Fourier transform mass spectrometers, coupled with advanced adaptive processing algorithms for data extraction, allow the detection and separation of over 10,000 chemicals in plasma. This includes
metabolites from 146 of the 154 pathways in the Kyoto Encyclopedia of Genes and Genomes (KEGG) human metabolic pathways,\textsuperscript{13} providing an effective coverage of central metabolic pathways. Importantly, greater than 90% of the KEGG metabolites have distinct elemental composition\textsuperscript{13} so that they can be measured based upon the accurate mass and mass resolution without requiring ion dissociation.\textsuperscript{14–16}

Metabolomic analysis of serum and plasma has revealed panels of metabolites that distinguish patients with cardiovascular disease,\textsuperscript{17} breast cancer,\textsuperscript{18,19} Parkinson disease,\textsuperscript{20} and diabetes\textsuperscript{21} from control patients. This technique is able to differentiate between individuals despite intraindividual variation due to factors such as dietary intake.\textsuperscript{14} We have established this metabolomic method as a powerful tool for assessing risk of ocular disease through identifying metabolites that discriminate neovascular AMD patients from controls.\textsuperscript{22}

The purpose of this study was 2-fold: first, to determine whether plasma metabolic profiling can detect systemic metabolic differences between POAG patients and controls; and second, to identify metabolites associated with POAG. Metabolic profiles incorporate both individual biomarkers and entire pathways associated with disease, providing a better understanding of POAG pathophysiology.

**METHODS**

**Ethics Statement**

This case control study was approved by the Vanderbilt University Human Research Protection Program. Research adhered to the tenets of the Declaration of Helsinki and was conducted in accordance with Health Insurance Portability and Accountability Act regulations. Written informed consent was obtained from all participants prior to study enrollment.

**Study Participants**

A total of 144 individuals were recruited from the Glaucoma Service at the Vanderbilt Eye Institute (VEI). Cases (n = 72) were diagnosed with POAG by a fellowship-trained glaucoma specialist (RWK). Diagnosis of POAG was based on optic nerve appearance characteristic of glaucoma, such as progressive cupping, thinning of the neuroretinal rim, notch formation, and disc hemorrhage. All cases had open iridocorneal angles as determined by gonioscopy examination. Patients with POAG underwent ancillary tests as indicated by their clinical findings, such as visual field testing (Humphrey visual field analyzer; Carl Zeiss Meditec, Dublin, CA, USA); fundus photography of the optic nerves (Zeiss 450+ fundus camera; Carl Zeiss Meditec); and retinal nerve fiber layer assessment with spectral domain optical coherence tomography (Cirrus OCT; Carl Zeiss Meditec). Although elevated intraocular pressure (IOP) was not included as a diagnostic criterion, IOP and central corneal thickness (CCT) were recorded for all POAG patients. Glaucoma treatment history was documented for cases.

Controls (n = 72) had no clinical signs of POAG, as determined by the same glaucoma specialist (RWK) using standardized diagnostic criteria, such as normal optic nerve and normal visual field. Exclusion criteria for cases included current or past retinal disease other than glaucoma, such as myopic retinopathy, diabetic retinopathy, or inherited retinopathy. Patients with optic nerve degeneration not secondary to glaucoma, active uveitis or ocular infection, or any ocular surgery within the 60 days prior to enrollment were also excluded. Smoking history was obtained from all participants; history and treatment of comorbidities were documented from each participant’s electronic medical record.

**Sample Collection**

At the time of study enrollment, blood was drawn from study participants using a 21- or 23-gauge butterfly needle into two 3-mL blood collection tubes, each containing 56 USP U lithium heparin. These tubes were centrifuged for 10 minutes at 4°C to pellet blood cells, and 2 mL supernatant were transferred to 1.5 mL conical tubes in 100 µL aliquots and immediately stored frozen at −80°C. Samples were subsequently shipped on dry ice to Emory University. Samples were not thawed, remained frozen during transit, and were frozen at −80°C at Emory prior to metabolomic analysis.

**Metabolomic Analysis**

Frozen plasma samples from all participants were thawed and analyzed by LC-MS at Emory University as previously described.\textsuperscript{14,15,22,23} For analysis, samples were batch randomized into 20-sample batches, with each batch containing an equal number of cases and controls in random order. Pooled reference plasma was run prior to and after each batch for quality control and quality assurance. Plasma sample aliquots (65 µL) were treated with 130 µL acetonitrile (2:1 vol/vol) containing an internal isotopic standard mix (3.5 µL per sample) as previously described.\textsuperscript{15,23} The internal standard mix for quality control (Supplementary Table S1) consisted of 14 stable isotopic chemicals covering a broad range of chemical properties represented in small molecules.\textsuperscript{15} Samples were mixed and placed on ice for 30 minutes prior to centrifugation for 10 minutes (16,100g at 4°C) to remove protein. The supernatants (10 µL) were loaded onto an autosampler (Accela Open Autosampler; Thermo Fisher Scientific, Inc., San Jose, CA, USA) maintained at 4°C and analyzed in triplicate using a mass spectrometer (Thermo LTQ Velos Orbitrap; Thermo Fisher Scientific, Inc.) and C18 chromatography (2.1 × 10 cm; Higgins Analytical, Inc., Targa, Mountain View, CA, USA). Elution was obtained with a formic acid/acetonitrile gradient at a flow rate of 0.35 mL/min for the initial 6 minutes and 0.5 mL/min for the remaining 4 minutes. The first 2-minute period consisted of a 5% solution A (2% [vol/vol] formic acid in water), 60% water, 35% acetonitrile, followed by a 4-minute linear gradient to a 5% solution A, 0% water, 95% acetonitrile. The final 4-minute period was maintained at 5% solution A, 95% acetonitrile. The mass spectrometer was set to collect from mass/charge ratio (m/z) 85 to 2000 over 10 minutes. Electro-spray ionization was used in positive mode.\textsuperscript{14,15} Optimal operating conditions determined for the mass spectrometer (Thermo Fisher Scientific, Inc.) are listed in Supplementary Table S2.

An adaptive processing software package (aplCMS, in the public domain at http://web1.sph.emory.edu/aplCMS/) designed for use with LC-FTMS data was used for peak extraction and quantification of ion intensities.\textsuperscript{24} This software provided m/z feature tables containing m/z values, retention time, and integrated ion intensity for each m/z feature, obtained through five major processing steps: (1) noise filter, (2) peak identification, (3) retention time correction, (4) m/z peak alignment across multiple spectra, and (5) reanalysis to capture peaks originally missed because of weak signal relative to the signal to noise filter.\textsuperscript{24} We used xMSanalyzer (in the public domain at http://sourceforge.net/projects/xmsanalyzer/) to enhance the feature detection process by performing systematic data re-extraction, statistical filtering, and data merger to enhance quality of data extraction.\textsuperscript{25} To ensure quality control, each sample was analyzed in triplicate, and a coefficient of variation (CV) was calculated for each m/z feature in each sample. Median CV, percent missing values, and a combined quality score were determined for each m/z.
Discriminatory m/z features were annotated on a targeted basis using the METLIN (in the public domain at http://metlin.scripps.edu/) and Kyoto Encyclopedia of Genes and Genomes (KEGG; in the public domain at http://www.genome.jp/kegg/) databases. The following adducts were used for annotation at ±10 ppm mass tolerance: M+H, M+Na, M+2H2O, M+H3O+, and M+2NaH.

**Ion Dissociation Mass Spectroscopy (LC-MS/MS)**

Ion dissociation mass spectrometry was carried out for representative DEM within the National Institute of Standards and Technology (NIST) Standard Reference Materials (SRM) 1950. Samples were analyzed using a high-resolution mass spectrometer (Q-Exactive; Thermo Fisher Scientific, Inc.) operated in positive ion mode and standard source conditions utilizing for the untargeted metabolic profiling. Analyte separation was accomplished on a C18 column and high performance liquid chromatograph (UltiMate 3000; Thermo Fisher Scientific, Inc.), with gradients optimized to match those used on the Surveyor system interfaced to the Velos Orbitrap HRMS. Prior to analysis, plasma proteins were precipitated using acetonitrile (2:1 vol/vol) and allowed to equilibrate for 30 minutes. Collision-induced ion dissociation was accomplished using high purity N2 at a normalized collision energy of 35%. Confirmation of the identities was completed by comparing the acquired spectra to online databases (MassBank, METLIN), in-silico fragmentation generators (MetFrag), and structural features.

**Data Analysis**

Descriptive statistics for demographic and clinical variables were calculated. Comparisons between cases and controls were made using the two-tailed t test for continuous data (e.g., age) and the two-tailed Fisher exact test for categorical data (e.g., sex, race, smoking, and comorbidities).

In order to increase confidence for selection of discriminating metabolites, xMSanalyzer was used to filter the data to include only m/z features for which at least 70% of either group had values. xMSanalyzer also provided batch normalization using ComBat to control for batch effects. Data then underwent log2 transformation to reduce heteroscedasticity and normalize results. We used 10-fold cross-validation accuracy criteria to minimize false positives and to determine the optimal log2 fold-change threshold for filtering, which was 0.05. To determine differences between cases and controls in the filtered m/z features, P values from LIMMA-based two-class comparison were adjusted for multiple hypothesis testing using the Benjamini-Hochberg false discovery rate (FDR) correction method. Two-way hierarchical cluster analysis (HCA) was used to visualize the relationship between participants and metabolites found to be different in POAG patients from controls. Analyses were conducted in R (in the public domain at http://www.r-project.org).

**Correlation Analyses**

Metabolome-wide Spearman correlation analysis was applied to the significant m/z features in order to capture any strongly correlated m/z features from the raw data. Associations with absolute Spearman correlation greater than 0.3 and FDR < 0.2 were included in the pathway analysis.

**Pathway Analysis**

Pathway analysis of metabolic features was completed using MetaboAnalyst (in the public domain at www.metaboanalyst.ca), a tool designed specifically for metabolomics studies. This software uses pathway enrichment analysis and pathway topology analysis to translate metabolic trends into defined pathways relevant to the study at hand. We used KEGG ID compound to cross-reference the KEGG metabolic pathway library.

**Data Access**

Data files relevant to this publication will be made available for nonprofit use by researchers upon request.

**RESULTS**

**Subject Characteristics**

The study population consisted of 144 patients, including 72 POAG cases and 72 controls. Comparisons between POAG patients and controls in terms of demographic and environmental data, as well as comorbid medical conditions, are presented in Table 1. The mean age of POAG patients did not significantly differ from that of controls, nor did the percentages of Caucasians, females, or smokers. There was no significant difference between cases and controls regarding coronary artery disease, hypertension, hyperlipidemia, diabetes, renal disease, or thyroid disease (Table 1). Disease-related clinical characteristics of POAG patients are presented in Table 2. General ocular health and characteristics are compared between POAG patients and controls in Supplementary Table S3. A higher percentage of POAG patients had best corrected visual acuity worse than 20/40 (P = 0.003). Control patients were less likely to have had cataract surgery (P = 0.034), but there was no difference between the groups when considering all types of ocular surgery (P = 0.352). Supplementary Table S4 shows comparisons between POAG patients and controls in...
terms of systemic medications. There was no difference between the groups in the percentage of patients taking any of nine types of systemic medications.

**High-Resolution Metabolomics Data**

High-resolution metabolomics uses LC-MS to detect chemicals as ions in the gas phase. The data extraction algorithms are designed to efficiently detect reproducible signals, most of which have not been studied sufficiently to know the identity of the associated chemical. Because of the accurate mass and mass resolution, however, the combination of the m/z of an ion and the retention time (RT) from the LC provides a largely unambiguous label even though the chemical identity may be unknown. Mass spectral data from the C18 LC-MS analysis of the 144 samples yielded 19,744 ions (defined by m/z and RT), each with associated ion intensity. Given the large number of features, we filtered the data as described in the Methods section to enhance the ability to see the major metabolic differences between POAG patients and controls. After filtering, 2440 m/z features were used for statistical analyses.

A metabolome-wide association study (MWAS) of POAG was performed by testing each of the m/z features for significant difference between the 72 POAG patients and 72 controls. The results of the analysis are visualized as a Manhattan plot (−log P versus RT; Fig. 1A). Since the analyses were performed using a reverse phase C18 column, hydrophobic chemicals interact more strongly than hydrophilic chemicals, typically resulting in longer retention time for hydrophobic chemicals. The results showed that most of the significant m/z features were chemicals with relatively hydrophobic characteristics, as would be expected from lipids. Because the m/z revealed that the chemicals were < 500 Da, the results indicated that the most significant metabolic differences were likely to include relatively simple lipids rather than larger, complex glycosphingolipids.

**Two-Way Hierarchical Cluster Analysis**

A two-way HCA was performed in order to gain a better understanding of how well the 41 m/z features discriminated POAG from controls and to test for similarity of distribution of the features among individuals. The results (Fig. 2) showed separation into two major clusters of individuals, with one consisting of 98% controls (51 individuals designated as group A), and the other consisting of 76% POAG patients (93 individuals designated as group B). As discussed below, group B appeared to have metabolic subgroupings, labeled as B1 (n = 63) and B2 (n = 30) in Figure 2. All three clusters, A, B1, and B2, became the focus of further analyses.

The 41 m/z features grouped into eight major clusters, with four clusters (3, 4, 5, and 7) containing metabolites with higher intensities in POAG patients than in controls and four clusters (1, 2, 6, and 8) containing metabolites with lower intensities in POAG patients (Supplementary Table S6). Combined, clusters 4, 5, and 6 contained the majority of the DEM (30 m/z features), demonstrating large groups of associated metabolites (Fig. 2). These results suggest that metabolic pathway and/or

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**Figure 1**. Metabolome-wide association study of POAG. (A) Type 1 Manhattan plot of metabolic features of POAG patients (n = 72) compared with controls (n = 72) shows that 41 of the 2440 m/z features differed at FDR = 0.05 (dotted horizontal line). Most of these correspond to chemicals with mass less than 500, suggesting that they are relatively small molecules, such as metabolic intermediates, peptides and/or relatively simple lipids. (B) Type 2 Manhattan plot shows that the significant features mostly eluted relatively late on the C18 chromatography analysis, indicating that the chemicals are likely to include lipids, such as steroids, terpenoids, fatty acids, and glycerolipids.
network differences exist between POAG patients and controls that could provide insight into underlying mechanisms or pathologic responses.

Annotation of Differentially Expressed Metabolites

Cluster 4 contained 12 ions that were higher in POAG than controls; 11 out of 12 eluted with retention times consistent with hydrophobic chemicals, suggesting lipid properties. Two had accurate m/z matches (m/z 422.3271, 232 seconds; 422.3245, 370 seconds) to palmitoylcarnitine; ion dissociation mass spectroscopy (LC-MS/MS) confirmed that the ion eluting at 370 seconds was indeed palmitoylcarnitine (Supplementary Fig. S1), while the other could be an isobaric species (monoglycerides, triglycerides, or cholane tetrols; Table 3). One group of cluster 4 ions (m/z 288.2884, 289.2918, 290.2953, 316.3200, and 317.3230) all putatively matched to sphingoid compounds. The 288.2884 m/z matched to C17 sphinganine, and two others (289.2918 and 290.2953) respectively matched the same chemical with one and two $^{13}$C atoms (increased by 1.0034). The 316.3200 m/z was 28.0316 atomic mass units greater than 288.2884, consistent with being a related lipid differing by two methylene ($-\text{CH}_2-$) units (i.e., C19-sphinganine). Additionally, m/z 286.2728 matched to both C17 sphingosine and heptadecanediol; although we attempted to obtain MS/MS data to confirm its identity, this feature was present at too low an abundance to obtain useful data. Highly correlated to this feature, m/z 244.2626 and m/z 272.2938 putatively matched pentadecanone and heptadecanone, respectively. These C15 and C17 chemicals are found in cereal grains. Of the remaining features in cluster 4, m/z 314.3041 matched heptadecanediol and m/z 292.2108 matched hydroxytetradecanedioic acid.

Cluster 5 contained 12 ions in higher abundance in group B1 than in group A or B2 (Fig. 2). Some of the ions were highly correlated, but there were no obvious multiple forms of single chemicals. The METLIN data included a variety of matches related to vitamin D (Table 3). Four putative matches to fungal sterols (vitamin D2) were present: ergosterol, m/z 438.3773, 531 seconds; ergostanol, m/z 466.4089, 531 seconds; hydroxyergocalciferol, m/z 377.3247, 532 seconds; and dihydroergocalciferol, m/z 443.3320, 530 seconds. Ion dissociation (MS/MS) study of m/z 377.3247 was consistent with the fragmentation pattern of hydroxyergocalciferol. Putative matches to vitamin D3 metabolites, which are not fungal sterols, were also present in cluster 5, including azacholecalciferol (m/z 350.3247, 533 seconds); and fluorohydroxycholecalciferol (m/z 399.3066, 533 seconds). These latter two ions

![Figure 2](https://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/934219/)
Table 3. Database Matches for Largest Three Clusters of Metabolites

<table>
<thead>
<tr>
<th>Cluster</th>
<th>P Value</th>
<th>m/z</th>
<th>RT, s</th>
<th>Presence in POAG</th>
<th>Match/Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>3.69E-10</td>
<td>422.2545</td>
<td>370</td>
<td>Higher</td>
<td>Palmitoylcarnitine</td>
</tr>
<tr>
<td>4</td>
<td>0.000133</td>
<td>422.2571</td>
<td>232</td>
<td>Higher</td>
<td>Monoglyceride (22:5); triglyceride (12:0/18:4/22:6), choline tetrol</td>
</tr>
<tr>
<td>4</td>
<td>2.37E-15</td>
<td>288.2884</td>
<td>345</td>
<td>Higher</td>
<td>C17 sphinganine</td>
</tr>
<tr>
<td>4</td>
<td>1.34E-14</td>
<td>289.2918</td>
<td>345</td>
<td>Higher</td>
<td>Probable 13C form of 288.280</td>
</tr>
<tr>
<td>4</td>
<td>2.95E-17</td>
<td>316.2490</td>
<td>352</td>
<td>Higher</td>
<td>Possible 288.2884 plus 2(C17)2, C19 sphinganine</td>
</tr>
<tr>
<td>5</td>
<td>6.62E-11</td>
<td>317.2520</td>
<td>368</td>
<td>Higher</td>
<td>Probable 13C form of 316.2520</td>
</tr>
<tr>
<td>4</td>
<td>3.06E-14</td>
<td>286.2728</td>
<td>350</td>
<td>Higher</td>
<td>C17 sphingosine; heptadecanediol</td>
</tr>
<tr>
<td>5</td>
<td>2.23E-08</td>
<td>244.2626</td>
<td>352</td>
<td>Lower</td>
<td>Pentadecanone</td>
</tr>
<tr>
<td>5</td>
<td>1.60E-19</td>
<td>272.2938</td>
<td>370</td>
<td>Higher</td>
<td>Hepatadecanone</td>
</tr>
<tr>
<td>4</td>
<td>7.63E-06</td>
<td>314.5041</td>
<td>370</td>
<td>Lower</td>
<td>Heptadecanediol</td>
</tr>
<tr>
<td>5</td>
<td>6.21E-07</td>
<td>292.2108</td>
<td>22</td>
<td>Higher</td>
<td>Hydroxytetradecanediolic acid</td>
</tr>
<tr>
<td>5</td>
<td>6.49E-12</td>
<td>399.3066</td>
<td>533</td>
<td>Higher</td>
<td>FluorohydroxyD3; (triterpene); DG(24/0/22:6)</td>
</tr>
<tr>
<td>5</td>
<td>4.96E-07</td>
<td>378.5564</td>
<td>518</td>
<td>Higher</td>
<td>No match</td>
</tr>
<tr>
<td>5</td>
<td>2.84E-10</td>
<td>333.2988</td>
<td>531</td>
<td>Higher</td>
<td>HODE-cholesterol ester</td>
</tr>
<tr>
<td>5</td>
<td>5.23E-10</td>
<td>377.3247</td>
<td>532</td>
<td>Higher</td>
<td>Hydroxyergocalciferol; DG(24/0/22:6)</td>
</tr>
<tr>
<td>5</td>
<td>1.25E-07</td>
<td>422.3820</td>
<td>529</td>
<td>Higher</td>
<td>Diapo-zeta-carotene (terpene)</td>
</tr>
<tr>
<td>5</td>
<td>7.48E-09</td>
<td>443.3320</td>
<td>530</td>
<td>Higher</td>
<td>Dihydroergocalciferol; cholestanediol</td>
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<td>5</td>
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<td>466.0898</td>
<td>531</td>
<td>Higher</td>
<td>Azacholecalciferol; decaprenol (terpene)</td>
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<td>5</td>
<td>4.45E-06</td>
<td>289.2726</td>
<td>531</td>
<td>Higher</td>
<td>Ergostanol</td>
</tr>
<tr>
<td>5</td>
<td>8.14E-12</td>
<td>366.3535</td>
<td>575</td>
<td>Higher</td>
<td>No match</td>
</tr>
<tr>
<td>5</td>
<td>7.75E-04</td>
<td>511.2540</td>
<td>534</td>
<td>Higher</td>
<td>HODE-methyl ester; monoglyceride</td>
</tr>
<tr>
<td>5</td>
<td>3.99E-04</td>
<td>433.7773</td>
<td>531</td>
<td>Higher</td>
<td>Ergosterol</td>
</tr>
<tr>
<td>5</td>
<td>3.26E-17</td>
<td>421.3158</td>
<td>394</td>
<td>Lower</td>
<td>Multiple vitamin D analogs; TG (12:0/16:0/20:5)</td>
</tr>
<tr>
<td>6</td>
<td>4.48E-22</td>
<td>422.3200</td>
<td>399</td>
<td>Lower</td>
<td>Probable 13C form of 421.3158</td>
</tr>
<tr>
<td>6</td>
<td>1.94E-13</td>
<td>376.2584</td>
<td>399</td>
<td>Lower</td>
<td>C19 sphingosine-1-phosphate</td>
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<tr>
<td>6</td>
<td>2.67E-08</td>
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<td>399</td>
<td>Lower</td>
<td>Probable 13C form of 376.2584</td>
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<tr>
<td>6</td>
<td>0.000255</td>
<td>482.4182</td>
<td>566</td>
<td>Lower</td>
<td>MG(24:1); TG (18/0/18/0/20:0)</td>
</tr>
<tr>
<td>6</td>
<td>0.000796</td>
<td>1256.1453</td>
<td>325</td>
<td>Lower</td>
<td>Complex glycosphingolipid</td>
</tr>
</tbody>
</table>

From HCA of features that differed between POAG patients and controls at FDR = 0.05. Searches were conducted at 10 ppm using METLIN. Many features had multiple isobaric matches and representative matches listed will require independent confirmation.

* Clusters are based on HCA as depicted in Figure 2.

also matched terpenes, a large class of compounds that contribute to the synthesis of mitochondrial ubiquinol, cholesterol, steroid hormones, and lipid-soluble vitamins. An additional terpene match (m/z 422.3820, 529 seconds) was also present. Among the other matches, m/z 286.2728, 575 seconds is noteworthy because it matched heptadecanediol, a metabolite from cereal grains related to metabolite matches for pentadecanone, heptadecanone, heptadecanediol, and heptadecanediol in cluster 4. Cluster 6 consisted of six m/z features that were quantitatively decreased in POAG relative to controls (Table 3). Two of the features appeared to be 13C forms of others. Assuming that these are 13C forms, three of the features matched triglycerides and two of the features matched C19 sphingosine-1-phosphate.

A small number of matches were present for the other clusters. Of these, cluster 8 contained two features which were decreased in POAG patients and matched MIPC and M(IP)2C, complex sphingolipids. Cluster 7 included a feature (m/z 366.3353, 575 seconds) that was higher in POAG patients and matched triglycerides and two of the features matched C19 sphingosine-1-phosphate.

To test whether pathway differences between POAG and controls could be identified with MetaboAnalyst, the broader network structure including the 41 DEMs and their correlated features was used. With a correlation threshold of |r| > 0.4 and FDR 0.05, palmitoylcarnitine (422.3245, 570 seconds) and C17-sphingosine/heptadecanediol (m/z 286.2728) from cluster 4 were closely associated with C19 sphingosine-1-phosphate from cluster 6 (Fig. 3). All three of these features were associated with hydroergocalciferol (m/z 377.3247) and the terpene (m/z 422.3280, 529 seconds) from cluster 5. Note that m/z 286.2728, which matches both C17-sphingosine and heptadecanediol, was negatively associated with C19-sphingosine-1-phosphate. These results show that the metabolites discriminating POAG from controls are present within a broad network structure involving interacting modules of metabolites.

**Pathway Analysis Using MetaboAnalyst**

To test whether pathway differences between POAG and controls could be identified with MetaboAnalyst, the broader network structure including the 41 DEMs and their correlated features was used. Three pathways were significant: galactose metabolism, fructose and mannose metabolism, and steroid hormone biosynthesis pathways (Table 4). Significant differences for steroid hormone biosynthesis (P = 0.00109) are consistent with the differences observed for terpenes and steroids, many of which are present as multiple isobaric species. MetaboAnalyst uses raw P value selection criteria

Correlation Among m/z Features

Metabolome-wide Spearman correlation analysis was applied to the 41 m/z features and produced a correlation network that included the DEM and their top 100 correlated features from the raw data. Like the HCA results, Spearman correlation demonstrated extensive relationships among the 41 m/z themselves. Using these data, we examined the relationships of clusters 4, 5, and 6 and by selecting representative m/z from each and testing for correlations with all other m/z features. With a correlation threshold of |r| > 0.4 and FDR 0.05, palmitoylcarnitine (422.3245, 570 seconds) and C17-sphingosine/heptadecanediol (m/z 286.2728) from cluster 4 were closely associated with C19 sphingosine-1-phosphate from cluster 6 (Fig. 3). All three of these features were associated with hydroergocalciferol (m/z 377.3247) and the terpene (m/z 422.3280, 529 seconds) from cluster 5. Note that m/z 286.2728, which matches both C17-sphingosine and heptadecanediol, was negatively associated with C19-sphingosine-1-phosphate. These results show that the metabolites discriminating POAG from controls are present within a broad network structure involving interacting modules of metabolites.

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followed by pathway enrichment analysis to test for pathway significance so that the differences in galactose metabolism \((P = 0.00066)\) and fructose and mannose metabolism \((P = 0.0015)\) may reflect changes in metabolites that were not significant after false discovery rate correction.

**DISCUSSION**

This metabolome-wide association study demonstrates the ability to identify systemic alterations in POAG patients that may provide new insight into disease etiology and pathological response. The most apparent trends involve palmitoylcarnitine, sphingolipids, vitamin D-related metabolites, and terpenes. The matches also reveal possible differences in odd-chain fatty acid metabolites and dietary lipids associated with cereal grains. These results support novel hypotheses for future targeted studies investigating causal mechanisms.

Numerous matches indicate changes in fatty acid metabolism in the POAG patients. Palmitoylcarnitine, which functions in the carnitine shuttle system for transport of fatty acids for β-oxidation in the mitochondria, was elevated in POAG cases, consistent with a previous study linking carnitine-palmitoyl transferase II deficiency to normal tension glaucoma. However, other factors can affect mitochondrial palmitoylcarnitine metabolism, suggesting that the increase in palmitoylcarnitine in POAG patients may reflect more generalized impairment of mitochondrial function rather than a specific characteristic of POAG. A previous study has found reduced levels of the omega 3 fatty acids docosahexaenoic (DHA) and eicosapentaenoic acid (EPA) in the blood of POAG patients, the former having even been tested as a treatment option for glaucomatous optic neuropathy. Putative matches to diacylglycerols and triacylglycerols containing DHA and EPA were elevated in POAG cases (Table 3).

Metabolites matching sphingosine and sphinganine were higher in POAG cases than in controls while sphingosine-1-phosphate (SIP) was lower; SIP is a fibrotic mediator and messenger also related to fatty acid metabolism. However, these \(m/z\) features putatively matched to sphingosine metabolites containing an odd number of carbons in their sphingoid backbones. Sphingosine metabolites in mammals typically contain backbones with an even number of carbons. Never-

![FIGURE 3. A metabolome-wide association study analysis of five selected \(m/z\) features (Y1 to Y5) with all other \(m/z\) features in the feature table. Correlation threshold \(|r| > 0.4\) at FDR 0.05.](https://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/934219/)

**TABLE 4. Metabolic Pathways Found by Metaboanalyst to Associate With POAG Status**

<table>
<thead>
<tr>
<th>Pathway</th>
<th>Total</th>
<th>Expected</th>
<th>Hits</th>
<th>Raw (P)</th>
<th>FDR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Galactose metabolism</td>
<td>41</td>
<td>1.10</td>
<td>6</td>
<td>0.0006577</td>
<td>0.041275</td>
</tr>
<tr>
<td>Steroid hormone biosynthesis</td>
<td>99</td>
<td>2.67</td>
<td>9</td>
<td>0.0010985</td>
<td>0.041275</td>
</tr>
<tr>
<td>Fructose and mannose metabolism</td>
<td>48</td>
<td>1.29</td>
<td>6</td>
<td>0.0015478</td>
<td>0.041275</td>
</tr>
</tbody>
</table>
theless, odd-numbered carbon chains are found in mammals, thought to be the result of alkylation or branching, and of dietary intake. Vitamin B12 finalizes the catabolism of odd-chain fatty acids, and accumulation of odd-chain fatty acid metabolites have also been noted to occur in some mammals due to vitamin B12 deficiency. This suggests a need for evaluation of multiple mechanisms of fatty acid metabolism in relation to glaucoma status.

A relatively large number of m/z putatively matched to vitamin D-related metabolites; 25-Azovitamin D3 is relevant as an inhibitor of the liver 25-hydroxylation of vitamin D3, potentially contributing to deficiency. Vitamin D deficiency is a potential risk factor for open angle glaucoma, and the interrelated metabolites expressed in this study may provide insight into this association. For example, sphingosine kinase (SphK1) is activated by 1α,25-dihydroxyvitamin D3, so inhibition of formation of 1α,25-dihydroxyvitamin D3 can potentially impact conversion of sphingosine to S1P and contribute to the overall metabolic signature of POAG patients. In principle, vitamin D2, a fungal sterol, could also interfere with these mechanisms. Importantly, no significant difference in the percentages of POAG patients and controls taking vitamin D supplements was found in this study (P = 0.317), making supplementation an unlikely explanation of the observed differences in vitamin D metabolites. Despite this, the difference between cluster B1 and B2 individuals (Fig. 3) appears to largely be explained by the sterol matches in metabolite cluster 5, suggesting a potential role for this association.

The interactions of these diverse metabolic systems are indicated by correlations of metabolites in Figure 3. The lipids function in energy metabolism and homeostasis, as well as in cell signaling. For instance, vitamin D can control formation of S1P as described above, but S1P has also been shown to function as a messenger to control steroid hormone biosynthesis. The detection of differences in metabolites matched to terpenes emphasizes this point. Terpenes are precursors to vitamin D, mitochondrial ubiquinol, cholesterol, and steroid hormones. However, they are also derived from diet so that perturbations could occur from diet or endogenous metabolism. The presence of multiple yeast and fungal sterols further indicates that complex interactions may contribute to POAG.

Finally, although no evidence for a causal relationship exists for the multiple cereal lipids that are increased in patients with POAG, the association warrants further investigation because diet or microbiome could impact these levels.

The detection of large numbers of unidentified chemical associated with POAG emphasizes strengths and limitations of this approach. The untargeted metabolomics design allows the evaluation of a wide range of metabolites in POAG patients and does not limit the analysis to a particular type of metabolic or known pathway. At the same time, the use of a single, standardized extraction protocol may have limited the extraction of certain specific metabolites, such as highly nonpolar lipids. Additionally, blood was drawn at patients’ regular eye clinic visits, making it likely that metabolites with a wider intraday variation than intersubject variation would not be identified in this study. Our standard sample extraction protocol for untargeted metabolomics is simple and reproducible, and does not call for the addition of antioxidants to plasma prior to storage at −80°C. Thus, some sample oxidation could occur even though samples are never thawed prior to analysis.

This study used high-resolution LC-MS with high mass resolution and mass accuracy, coupled with advanced adaptive processing algorithms for data extraction. Such techniques can accurately identify metabolites within 5 to 10 ppm. However, the putative matches to the metabolomics databases require validation by ion dissociation mass spectrometry (MS/MS). The population size is relatively small and from a limited geographical area, so that further testing in an independent replication cohort is warranted. Initial MS/MS analysis in this study did confirm the identity of two metabolites: palmitoyl-carnitine and hydroxyergocalciferol. Finally, although there were no major differences in comorbidities (Table 1) or systemic medications (Supplementary Table S4) between POAG patients and controls, the metabolic changes linked with POAG in this study could be the result of disease or treatment status, and causality cannot be inferred from this association study.

In summary, this MWAS of 72 POAG patients and 72 controls shows that systemic metabolic signatures differentiate cases and controls. The pathway and network analyses show that complex pathway interactions occur and provide a basis to formulate testable hypotheses concerning potential metabolic and dietary factors that could impact disease occurrence and progression. These results indicate that factors impacting mitochondrial metabolism of fatty acids, steroid biosynthesis, and sphingolipid metabolism are worthy of further study. The results also suggest that vitamin D2 and D3 may be mechanistically important. Overall, the results illustrate that metabolomics offers new avenues to explore the pathophysiology of POAG.

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