EFEMP1 rare variants cause juvenile-onset open angle glaucoma in families from the Philippines

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July 19, 2021

Abstract

Juvenile open angle glaucoma (JOAG) is a severe type of glaucoma with childhood onset and dominant inheritance. Using exome sequencing we identified 3 independent families from the Philippines with rare EFEMP1 variants (c.238A>T, p.Asn80Tyr; c.1480T>C p.Ter494Glnext*29; and c.1429C>T, p.Arg477Cysteine) co-segregating with disease. Affected variant carriers (N=34) exhibited severe disease with average age of onset of 16 years and with 76% developing blindness. To investigate functional effects, we transfected COS7 cells with vectors expressing the three novel EFEMP1 variants and showed that all three variants found in JOAG patients caused significant intracellular protein aggregation and retention compared to wild type and also compared to EFEMP1 variants associated with other ocular phenotypes including an early-onset form of macular degeneration, Malattia Leventinese/Doyne’s Honeycomb retinal dystrophy. These results suggest that rare EFEMP1 coding variants can cause JOAG through a mechanism involving protein aggregation and retention, and that the extent of intracellular retention correlates with disease phenotype. This is the first report of EFEMP1 variants causing JOAG, expanding the EFEMP1 disease spectrum. Our results suggest that EFEMP1 mutations appear to be a relatively common cause of JOAG in Filipino families, an ethnically diverse population.

Introduction

Glaucoma is one of the most heritable of all human diseases (Wang et al., 2017) and exhibits both Mendelian (childhood onset) and complex (adult onset) inheritance (Wiggs and Pasquale, 2017). Affected patients typically develop elevated intraocular pressure (IOP) that causes irreversible degeneration of retinal ganglion cells and the optic nerve (Weinreb et al., 2016). Increased IOP in most patients is caused by impaired drainage of intraocular fluid from the eye through the trabecular meshwork and outflow pathways (Weinreb et al., 2014).
Patients with childhood-onset glaucoma, defined as diagnosis before age 40, are affected by severe disease characterized by very high IOP, and without treatment are likely to become blind over their lifetimes (Papadopoulos et al. 2020). Juvenile-onset primary open-angle glaucoma (JOAG) is one type of childhood glaucoma that typically exhibits dominant inheritance. MYOC mutations account for disease in 15-20% of JOAG families primarily with European Caucasian ancestry and MYOC does not appear to be a major cause of disease in other ethnic groups (Liu et al., 2021). To investigate the genetic etiology of JOAG in an ethnically diverse population, we identified and recruited 14 JOAG pedigrees from regions throughout the Philippines.

EFEMP1 (EGF-containing fibulilike extracellular matrix protein 1) also known as Fibulin 3 is a member of the Fibulin family of extracellular matrix proteins characterized by tandem arrays of EGF (epidermal growth factor)-like domains and a C-terminal fibulin-type module (Figure 1B) (Kobayashi et al., 2007). Among the fibulin family members, EFEMP1 has higher ocular expression (Wagner et al., 2013) and is known to contribute to other ocular phenotypes. Common SNPs near EFEMP1 on chromosome 2p16 have been associated with adult-onset glaucoma (POAG) (Gharahkhani et al., 2021) as well as IOP (Khawaja et al., 2018) and cup-to-disc ratio (CDR) an optic nerve quantitative trait related to POAG risk (Springelkamp et al., 2015). A low frequency EFEMP1 variant p.Arg140Trp, present in 2 individuals in gnomAD has been identified in 5 affected individuals from an adult-onset POAG African-American family (Mackay et al., 2015) and a stop loss variant (Ter494Gluext*29) has been reported in 3 affected members of a Chinese POAG family (Liu et al., 2020). A single EFEMP1 missense allele (p.Arg345Trp) is known to cause Malattia Leventinese (MLVT) (also known as Doyne’s Honeycomb dystrophy (DHRD; MIM:126600), a rare autosomal dominant retinal degeneration characterized by extracellular deposits (drusen) beneath the retinal pigment epithelium (RPE) (Stone et al., 1999). Among the many patients affected by MLVT/DHRD, only p.Arg345Trp has been identified as a causative mutation. Recently it has been suggested that the Arg345 residue is a mutation hot spot and that founder effects are not likely to account for the predominance of this mutation in MLVT/DHRD patients (Vaclavik et al., 2020). Notably, wildtype EFEMP1 is a secreted protein, while the p.Arg345Trp mutations leads to protein misfolding and reduced secretion (Marmorstein et al., 2002).

Here, we report three independent families from the Philippines with novel EFEMP1 rare variants segregating with JOAG. Functional studies suggest that the disease-associated EFEMP1 variants cause protein misfolding and intracellular aggregation and retention. These results expand the spectrum of EFEMP1 associated disease to include childhood glaucoma.

Methods

Ethical approval for this study was obtained from the Institutional Review Board of the Manila Doctors Hospital and Cebu Velez General Hospital, Philippines and the Mass General Brigham (MGB) IRB. In accordance with the tenets of the Declaration of Helsinki, written informed consent was obtained from the study subjects after explanation of the nature and possible consequences of the study.

Study recruitment

Subjects with juvenile open angle glaucoma (JOAG) were recruited from ophthalmology clinics. When possible, subjects underwent an ophthalmologic exam, which included visual acuity assessment, slit-lamp examination to evaluate the anterior segment, dilated fundus examination to document the structural changes of glaucoma, and gonioscopic examination to evaluate the angle structure. JOAG was defined as: 1) absence of secondary causes of glaucoma; 2) IOP > 21 mmHg; 3) open anterior chamber angle on gonioscopy (grade 3 or 4 of modified Shaeffer classification); 4) characteristic optic disc damage and/or typical visual field loss; and 5) IOP >21 mmHg in at least one eye in any family member aged between 3 and 40 years old. Pedigree information and disease status were ascertained from probands and family members. Asymptomatic family members were also invited to participate in the study and undergo a clinical examination and blood draw. Venipuncture was done on each subject and 10 cc of venous whole blood was obtained. Genomic DNA was extracted from the blood samples with the DNeasy Blood DNA extraction kit (Qiagen, Valencia, CA) according to the manufacturer’s instructions.
Exome sequencing

Whole exome sequencing (WES) was performed at the Massachusetts Eye and Ear Ocular Genomics Institute using Agilent SureSelect Human All Exon v6 with the Illumina HiSeq 2000, or at the Broad Institute of MIT and Harvard using a TWIST Biosciences custom exome bait with the Illumina Novaseq 6000. We used the Ocular Genomics Institute bioinformatics pipeline (Consugar et al., 2015) for alignment and variant calling that includes the Burrows-Wheeler aligner (Li et al., 2010), GATK (https://gatk.broadinstitute.org/), GERP (http://mendel.stanford.edu/SidowLab/downloads/gerp/), Exome Aggregation Consortium (ExAC) (http://exac.broadinstitute.org), SIFT (http://sift.jcvi.org/), and PolyPhen-2 (http://genetics.bwh.harvard.edu/pph2/). A coverage depth cutoff of 10x was applied. Heterozygous was defined as a fraction of a variant base between 0.25–0.75 and homozygous was defined as above 0.75. Average coverage was 105x for 99% of coding sequences.

Variant filtering

Ten genes (MYOC, CYP1B1, LTBP2, PITX2, FOXC1, PAX6, TEK, ANGPT1, LMX1B, CPAMD8) known to cause various forms of early-onset glaucoma were initially screened for pathogenic mutations. If no plausible causative variants were identified, the exome data was examined for novel pathogenic variants. Disease-causing mutations are expected to be rare, disrupt protein function, and have high estimates of pathogenicity. Variants were filtered using 1) minor allele frequency (MAF) from the Exome Aggregation Consortium (ExAC) database (MAF less than 0.001 were retained), 2) tools to score variants according to phylogenetic conservation (GERP) and its effect on protein structure (SIFT, PolyPhen, conserved predicted pathogenic variants were obtained), 3) ocular expression, and any known functional or human disease involvement (ocular expression was prioritized) and 4) examining segregation in all available family members. The family A and B pedigree structures suggested dominant inheritance; hence, heterozygous variants were prioritized. Sanger sequencing was used to investigate segregation of candidate variants in all available family members.

EFEMP1 expression vector

To evaluate the cellular expression and localization of disease-associated EFEMP1 variants we used a pENTR/D-TOPO entry vector (Invitrogen, Carlsbad, CA) with cloned wild-type EFEMP1 cDNA (provided as a gift from Dr. Rosario Fernandez-Godino, (Garland et al., 2021). The EFEMP1 mutants (p.Asn80Tyr, p.Arg140Trp, p.Arg477Cys) were created using the QuikChange Site-Directed Mutagenesis kit (Agilent Technologies, Santa Clara, CA). To create the p.Ter494Glnext*29 mutation, a gBlocks Gene Fragment (Integrated DNA Technologies, Iowa) was designed and cloned in to a pENTR/D-TOPO entry vector. The wild-type and mutant entry clones were verified by sequencing analysis and moved into a Gateway destination expression vector modified to contain an N-terminal V5 epitope tag in-frame (pCAG-V5-IRESEGFP) (Zhang et al., 2012) by LR recombination. Plasmid DNA was purified using the EndoFree Plasmid Maxi kit (Qiagen, Valencia, CA).

Cell culture

COS-7 cells were used to express EFEMP1 protein. The COS-7 cells were seeded in 6-well plates at a concentration of 100,000 cells per mL and allowed to grow in DMEM + 10% FBS condition media. The cells were transfected with expression clones encoding wild-type or mutant EFEMP1 cDNA using Lipofectamine 3000 (Invitrogen, Carlsbad, CA) 24 hours post-seeding at 80-90% confluence.

Immunocytochemistry

To visualize the cellular localization of EFEMP1 protein, cells were seeded on clear coverslips (Neuvitro Corp., Camas, WA). In order to visualize the endoplasmic reticulum, CellLight ER-RFP, BacMam 2.0 (Invitrogen, Carlsbad, CA) was added to each well during transfection. The cells were processed for immunocytochemistry 48 hours after transfection. Condition media was aspirated, coverslips washed with PBS, and fixed in 4% PFA. Cells were permeabilized using 0.5% Triton X-100 and blocked in 3% BSA with PBS, followed by incubation in V5 Tag Monoclonal Antibody (Thermofisher, Waltham, MA) overnight. Coverslips
were retrieved and incubated in Goat anti-Mouse IgG (H+L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor Plus 647 (Invitrogen, Carlsbad, CA) for 2 hours and DAPI (Thermofisher, Waltham, MA) for 1 minute. Coverslips were mounted on clear slides with ProLong Glass Antifade Mountant (Invitrogen, Carlsbad, CA). Imaging of the transfected cells was done using a confocal laser scanning microscope (SP8, Leica Microsystems, Buffalo Grove, IL).

**Western blots**

To quantify transfected EFEMP1 protein, cells were harvested 48 hours post transfection (average transfection efficiency 75%) (Table S1). Whole cell lysates were prepared by careful aspiration of condition media and adding cold RIPA buffer (Sigma-Aldrich, St. Louis, MO) and Pierce Protease Inhibitor (Thermofisher, Waltham, MA) to each well and agitating for 30 minutes at 4°C. After incubation, cells were transferred to Eppendorf tubes and centrifuged for 30 minutes at 4°C. The supernatant was retrieved and the protein concentration was determined using the Pierce BCA Protein Assay Kit (Thermofisher, Waltham, MA). For each vector group, 5ug of protein was mixed with 4X Protein Sample Loading Buffer (Li-Cor Biosciences, Nebraska) and water for a total volume of 12ul. Each sample was denatured by heating at 95 degC for 3 minutes and loaded in to a 15 well 4-20% Mini-PROTEAN(r) TGX Precast Protein Gel (Bio-Rad, Hercules, CA). Electrophoresis was performed at 120v for approximately 1.5 hours. The gel was retrieved and transferred to a PVDF blotting membrane using the iBlot 2 system (Invitrogen, Carlsbad, CA). For normalization of protein loading, total protein staining was done using the Revert 700 Total Protein Stain Kit (Li-Cor Biosciences, Nebraska), following the manufacturer’s protocol. After imaging in the 700nm channel using the Odyssey(r) Imaging System (Li-Cor Biosciences, Nebraska), the membrane was blocked for 1 hour at room temperature using Intercept(r) Blocking Buffer (Li-Cor Biosciences, Nebraska), then overnight incubation with Anti-Fibulin-3 Antibody (mab3-5): sc-33722 (Santa Cruz Biotechnology, Dallas, TX) diluted with blocking buffer to a final concentration of 1:100. After overnight incubation, the membrane was incubated in IRDye(r) 800CW secondary goat anti-mouse antibody (Li-Cor Biosciences, Nebraska), at a final concentration of 1:10,000 for 1 hour and then imaged in the 800nm channel. Western blot quantification and analysis was carried out using ImageStudioLite(r) Software (Li-Cor Biosciences, Nebraska). Relative EFEMP1 abundance was calculated as the ratio between the EFEMP1 signal and the corresponding total protein signal for each sample. Three biological repeats were done for each EFEMP1 condition. Differences between mutant and wildtype EFEMP1 were tested using Student’s t-test with Bonferroni correction.

**Statistical Analysis**

All the analysis were done using GraphPad Prism software (GraphPad Software, San Diego, CA). Adjusted p value [?] 0.05 was considered significant.

**RESULTS**

**Identification of rare coding EFEMP1 variants in three families with JOAG.**

Whole exome sequencing was completed for selected affected and unaffected family members, and exome data was filtered to retain rare variants (MAF < 0.1%) that were protein altering and predicted to be pathogenic using in silico programs. Mutations were identified in known disease-causing genes in 4 families, three with MYOC mutations (p.Ter505Trpext*42; p.Gln337Arg; p.Thr438Ile) and one with a PAX6 mutation (p.Tyr296*). Novel variants of interest (Table S2) were further evaluated for co-segregation with disease in each pedigree using Sanger sequencing. Rare EFEMP1 (MIM: 601548) coding variants (GenBank: NM_001039348.3) were identified in all affected individuals of 3 of the remaining families as follows: (c.238A>T, p. Asn80Tyr) in 16 affected individuals of Family A and not in 18 unaffected family members older than age 18; (c.1480T>C, p. Ter494Glnext*29) was identified in 17 affected family members in family B and not in any family members older than age 18. One Family A member (age 13) and 4 Family B members (ages 7-12) were identified as mutation carriers, but are likely to be too young to manifest disease. Additionally, a third variant (c.1429C>T, p.Arg477Cys) was identified in a single case (Family C) (Figure 1A).

Patients with EFEMP1 variants (N= 34 in total for all three families) exhibited severe disease with average
age of disease onset of 16 years (range 3-43). Affected individuals had much higher than average IOP (28 mmHg) and 76% were blind in at least one eye (Table 1). The majority of patients required surgical treatment for IOP elevation, however not all patients had access to clinical care. Retinal examination did not reveal evidence of subretinal deposits (drusen) characteristic of MLVT/DHRD in affected individuals (Figure 2).

The three variants p.Asn80Tyr, p.Ter494Glnex*29, and p.Arg477Cys, identified in affected individuals are not present in population databases including gnomAD and TOPMed and have not been previously reported (the stop loss variant, p.Ter494Glnex*29 is not the same variant as reported in a small Chinese adult onset glaucoma family) (Liu et al., 2020). All three variants are predicted to be deleterious by in silico programs SIFT and Polyphen2 (Table S3). p.Asn80Tyr is located within the first EGF-domain, while p.Arg477Cys and p.Ter494Glnex*29 are within the terminal fibulin-like carboxy terminal domain. p.Ter494Glnex*29 results from replacement of the wild type stop codon with a glutamine residue and the addition of 29 amino acid residues to the polypeptide before a stop codon is encountered (Figure 1B).

Expression and localization of EFEMP1 wild-type and mutant proteins

EFEMP1 is expressed in multiple ocular anterior segment cell types especially the trabecular meshwork beam cells and juxtacanilicular matrix cells (Figure 3) (van Zyl et al., 2020), two structures necessary for outflow of intraocular fluid (aqueous humor) and maintenance of normal IOP levels. EFEMP1 ocular expression is similar to Myocilin (MYOC) also an extracellular matrix protein involved in JOAG. MYOC mutations cause protein misfolding and endoplasmic reticulum aggregation, and efforts to reduce mutation-related misfolding can result in lower IOP in animal models (Orwig et al., 2014; Jain et al., 2017).

To assess the functional effects of the JOAG-related EFEMP1 variants, we hypothesized that these variants could cause protein misfolding, aggregation and intracellular retention similar to that observed for the p.Arg345Trp MLVT/DHRD variant and for JOAG-related MYOC mutations. To test this hypothesis we transfected COS7 cells with Gateway destination expression vectors (pCAG-V5-IRES-EGFP) (Zhang et al., 2012) for wildtype EFEMP1 cDNA, the three JOAG-related EFEMP1 variants, and also the MLVT/DHRD variant (p.Arg345Trp) and the POAG-related variant (p.Arg140Trp). Imaging transfected cells showed that wild type EFEMP1 was distributed throughout the cell, while significant intracellular aggregation was observed for each JOAG-related variant (Figure 4). Co-localization with an endoplasmic reticulum (ER) marker, CellLight ER-RFP, BacMam 2.0 (Invitrogen, Carlsbad, CA), suggests that EFEMP1 protein aggregates are formed in the vicinity of the ER. In contrast, the p.Arg345Trp variant exhibited more diffuse cellular distribution as did the POAG related variant p.Arg140Trp.

To measure intracellular protein retention, we collected an equal number of transfected cells from experiments using the wild type vector, the 3 JOAG-related vectors and the p.Arg345Trp and p.Arg140Trp vectors. Using Western blot assays (Figure 5A, Figure S1) we determined the fraction of EFEMP1 protein retained within the cell in comparison to overall protein and relative to wild type protein (Figure 5B). Compared to wild type cells intracellular EFEMP1 protein was significantly increased for all three of the JOAG-related variants (P < 0.05). JOAG-related intracellular protein was also increased compared to the MLVT/DHRD variant p.Arg345Trp and the POAG-related p.Arg140Trp.

Discussion

Collectively, these data support the hypothesis that EFEMP1 coding variants can cause JOAG by a mechanism that involves intracellular protein aggregation and retention. Similarly, JOAG-causing MYOC mutations also promote formation of intracellular protein aggregates and the extent of protein aggregation appears to correlate with disease severity (Patterson-Orazem et al., 2019). Misfolded myocilin is hypothesized to cause ER stress leading to cellular dysfunction and potentially apoptosis (Yam et al., 2007), however the specific cells that are impacted and the underlying molecular events are not known. Studies of human and mouse MYOC knockouts (Kim et al., 2001; Pang et al., 2002; Wiggs et al., 2001) and mouse knockins (Kim et al., 2001; Zode et al., 2011) indicate that MYOC mutations are gain-of-function. The observation that the retina and optic nerves of EFEMP1 knockout mice are anatomically normal (McLaughlin et
al., 2007; Stanton et al., 2017; Daniel et al., 2020) suggests that \( EFEMP1 \) loss of function also does not underlie glaucoma development. \( MYOC \) and \( EFEMP1 \) are among a group of proteins expressed in ocular extracellular matrix that also includes Thrombospondin1 (\( THBS1 \)), and Angiopoietin-like 7 (\( ANGPTL7 \)) among other proteins with potential glaucoma involvement (Wirtz et al., 2021; Tanigawa et al., 2020). Although the specific mechanisms underlying the contribution of ECM proteins to glaucoma is not known, preventing mutant \( MYOC \) expression (Jain et al., 2017) or encouraging secretion of misfolded myocilin can reduce intraocular pressure in mice (Zode et al., 2011; Zode et al., 2021), and similar approaches may also be therapeutically useful for patients with \( EFEMP1 \) mutations.

In this study we showed that the JOAG associated variants demonstrate increased intracellular protein retention compared with the MLVT/DHRD variant p.Arg345Trp. \( EFEMP1 \) p.Arg345Trp has recently been shown to effect cholesterol efflux (Tsai et al., 2021) which may potentially impact the contribution of this mutation to the characteristic extracellular protein aggregation observed in this disease (Fu et al., 2007). While some POAG genomic loci include genes that may influence cholesterol efflux (\( ABCA1 \), \( ARHGEF12 \), \( CAV1/2 \)) (Gharahkhani et al., 2021; Springelkamp et al., 2015; Jacobo-Albavera et al., 2021; Wang et al., 2014; Okuhira et al., 2010), the role of cholesterol metabolism in glaucoma is not clear and given the involvement of \( EFEMP1 \) in JOAG could be interesting to explore further. It is also of interest that only the single missense allele p.Arg345Trp is known to cause MLVT/DHRB, yet at least 3 protein variants can cause glaucoma. It is possible that \( EFEMP1 \) missense alleles are more likely to effect cells of the ocular outflow pathway than the retinal pigment epithelial cells involved in MLVT/DHRB. Interestingly, patients with MLVT/DHRB are not known to be at increased risk of glaucoma, and the JOAG patients with \( EFEMP1 \) mutations do not have any evidence of the MLVT/DHRB retinal dystrophy (Figure 2).

Additionally, we show that the JOAG \( EFEMP1 \) variants demonstrate increased intracellular retention compared to a possible POAG associated variant, p.Arg140Trp found in a family with 5 members affected by adult-onset POAG (Mackay et al., 2015), a finding that may be related to the less severe and genetically complex adult-onset POAG. Our results could support a contribution of p.Arg140Trp to POAG in this family, which is interestingly of African American ancestry. Exome-based studies of African American POAG cases or families have not yet been completed and further investigation in this population could reveal additional \( EFEMP1 \) disease-related variants. A stop lost variant (Ter464Gluext*29) similar to the mutation affecting JOAG family B (Ter464Gluext*29) has recently been described in 3 members of a Chinese adult POAG family (Liu et al., 2020). While individuals in this family are described as affected by POAG, the age of disease onset is the mid-20s which would meet our diagnostic criteria for JOAG and is consistent with our results.

\( EFEMP1 \) is located on chromosome 2p16 within a genomic region that was initially identified as a POAG genomic locus (GLC1H) in a linkage study of a Jamaican family and European Caucasian families from the UK (Suriyapperuma et al., 2007). Subsequently linkage to this region has also been observed in four Chinese families, including two with disease onset before age 40 (Liu et al., 2012; Liu et al., 2008). POAG candidate association studies have also implicated SNPs in the 2p16 region in cases and controls from Barbados (Jiao et al., 2009), an African-American cohort (Liu et al., 2010), Chinese (Chen et al., 2012) and South Indian cohorts (Balasubbu et al., 2021). Together with our results, these studies support \( EFEMP1 \) as the GLC1H gene and also could suggest that \( EFEMP1 \) is more commonly associated with glaucoma in non-white populations.

Our findings also suggest a phenotype-genotype spectrum that correlates the extent of intracellular protein retention with disease phenotype and extend the \( EFEMP1 \) phenotypic spectrum to include severe childhood-onset glaucoma. The mutations causing JOAG cause more severe intracellular protein retention compared to the MLVT/DHRD \( EFEMP1 \) variant and also to a variant found in an adult-onset glaucoma family. Further study will be required to determine the mechanism underlying intracellular protein retention as well as the molecular events leading to elevated IOP and glaucoma.

This study investigates the etiology of JOAG in a collection of affected families from the Philippines, a population with diverse ancestry including Asian, African and European Caucasian origins (Larena et al.,
2021). Among the pedigrees that we have collected we find that \textit{EFEMP1} variation is a relatively common cause of childhood glaucoma (20%), equal to \textit{MYOC} in this population. Interestingly, \textit{EFEMP1} variants have not been observed in any prior studies of childhood glaucoma which have focused primarily of families with European Caucasian or Asian ancestry (Allen et al., 2015; Huang et al., 2018). As well, \textit{MYOC} and other currently known childhood gene mutations have infrequently been identified in patients with African ancestry (Liu et al., 2012). These results suggest that investigation of diverse populations such as this Filipino cohort will be necessary to develop a more comprehensive set of childhood glaucoma genes.

The discovery of genes causing childhood glaucoma makes it possible to use genetic testing to inform genetic counseling for affected families. Treatment initiated at early stages of disease can delay irreversible optic nerve degeneration and provide the best chance that an affected child will maintain useful sight throughout their lifetime. Informed genetic counseling makes it possible to create surveillance and treatment plans for mutation carriers and alleviates the burden of screening family members without disease-causing mutations. Currently however a molecular diagnosis can only be achieved for approximately 20-25% of cases based on known genes (Allen et al., 2015). Discovery of novel disease-causing genes such as \textit{EFEMP1} is needed to improve the overall diagnostic yield and effectiveness of childhood glaucoma genetic testing.

In conclusion we have identified 3 different \textit{EFEMP1} coding variants that segregate with a severe form of glaucoma affecting children in 3 independent families from the Philippines. Our results suggest that disease-associated variants cause significant intracellular EFEMP1 aggregation and retention and that the extent of intracellular retention appears to be correlated with EFEMP1-related disease phenotypes. This study further supports a role for EFEMP1 in ocular extracellular matrix and in regulation of intraocular fluid dynamics and IOP and provides new opportunities for genetic testing and therapeutic intervention.

Data availability statement

The authors confirm that the data supporting the findings of this study are available within the article or supplementary information. All variants identified in the present study have been submitted to ClinVar (https://www.ncbi.nlm.gov/clinvar/) and are awaiting accession numbers.

Acknowledgments

Supported by the ARVO Foundation David L. Epstein award (ERAC) and NIH/NEI R01 EY031830 ad P30EY014104 (JLW)

Declaration of Interests

JLW has received research support from Aerpio pharmaceuticals and is a consultant for Aerpio, Allergan, Maze, Editas, Regenxbio and Avelino.

Web Resources

gnomAD, https://gnomad.broadinstitute.org/

TOPMed, https://bravo.sph.unich.edu/freeze8/hg38/

SIFT, https://sift.bii.a-star.edu.sg/

Polyphen2, http://genetics.bwh.harvard.edu/pph2/


OMIM, https://www.omim.org/

References


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Mutation carrier, no clinical signs of glaucoma

Table 1. Clinical findings in subjects with *EFEMP1* rare coding variants. All clinical data listed were obtained during study recruitment. Normal IOP is less than 21 and visual acuities of 20/200 or greater meet the definition for blindness. NLP, LP, CF, and HM also meet the definition for blindness. Abbreviations: Single dash (-): exam not performed because of poor view, corneal opacification, or phthisis.
bulbi; na: not applicable; PAS: peripheral anterior synechiae; NLP: no light perception; LP: light perception; CF: counting fingers; HM: hand movement; M: male; F: female; OD: right eye; OS: left eye.

Figure titles and legends

**Figure 1A.** Family pedigrees with juvenile-onset open-angle glaucoma (JOAG) and **EFEMP1** mutation status. Arrow indicates the proband. Squares indicate male subjects; and circles refer to female subjects. Slash through indicates a deceased individual. Solid symbols indicate subjects diagnosed with JOAG; half-filled symbols indicate subjects who carry the **EFEMP1** variant but with no clinical signs of glaucoma; and open symbols refer to subjects with no glaucoma and no **EFEMP1** variant. Genotypes are heterozygous mutant (m/+) and wild type (+/+).

**Figure 1B.** Schematic diagram of **EFEMP1** and evolutionary conservation. **EFEMP1** Genbank transcript NM_001039348.3 and color-coded exons corresponding to the protein domain. The mutations p.Asn80Tyr, p.Arg477Cys, and p.Ter494Glnext*29 are located at highly conserved domains across several organisms.

**Figure 2.** Fundus images from a patient from Family B with **EFEMP1** variant p.Ter494Glnext*29 (B-V:5). The image show damage to the optic nerve but no evidence of subretinal deposits (drusen) characteristic of Malattia Leventinese/Doyne Honeycomb dystrophy.

**Figure 3.** Single cell RNA sequencing **EFEMP1** expression and known early onset glaucoma genes from the human aqueous humor outflow pathway. Comparative expression levels of **EFEMP1** and genes known to cause various types of early onset glaucoma. Plot generated from the Broad Institute of MIT and Harvard’s Single Cell Portal using data from “Cell atlas of aqueous humor outflow pathways in eyes of humans and four model species provides insight into glaucoma pathogenesis” by van Zyl et al, 2019.

**Figure 4.** Expression of **EFEMP1** in COS-7 cells. Cultured COS-7 cells transfected with wildtype and variant **EFEMP1** and processed with immunocytochemistry and the following stains: DAPI (nucleus), CellLight ER-RFP (endoplasmic reticulum), Alexa 647 (**EFEMP1** protein), GFP (GFP expression reporter). Imaged using Leica SP8 confocal microscope using 63x objective (glycerol immersion), digital zoom 2x.

**Figure 5.** **EFEMP1** expression in COS7 cells. 
A. Western blot of wild-type (WT) and mutant EFEMP1 proteins. Transfected COS7 cell-lysates showing expression of WT and mutant EFEMP1. Control group consist of cell-lysates from non-transfected COS7 cells. B. Ratio of intracellular EFEMP1 protein to total protein for wild type and each variant. Relative EFEMP1 abundance was calculated as the ratio between the EFEMP1 signal and the corresponding total protein signal in each sample measured using Revert 700 total protein stain. Three biological repeats were done for each EFEMP1 condition. Differences between mutant EFEMP1 and wildtype EFEMP1 were tested using Student’s t-test with Bonferroni correction. Error bars indicate the standard error. *p<0.05, **p<0.001.