Expression Profile of the Matricellular Protein Osteopontin in Primary Open-Angle Glaucoma and the Normal Human Eye

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PURPOSE. To characterize the role of osteopontin (OPN) in primary open-angle glaucoma (POAG) and normal eyes.

METHODS. OPN quantification was performed by enzyme-linked immunosorbent assay in aqueous humor (AH) obtained from human donor eyes (POAG and normal) and surgical samples (POAG and elective cataract removal). OPN expression and localization in whole eye tissue sections and primary normal human trabecular meshwork (NTM) cells were studied by Western blot and immunohistochemistry. Latanoprost-free acid (LFA)–treated NTM cells were analyzed for OPN gene and protein expression. Intraocular pressure was measured by tonometry, and central corneal thickness was measured by optical coherence tomography in young OPN+/− and wild-type mice.

RESULTS. OPN levels were significantly reduced in donor POAG AH compared with normal AH (0.54 ± 0.18 ng/μg [n = 8] vs. 0.77 ± 0.23 ng/μg [n = 9]; P = 0.039). A similar trend was observed in surgical AH (1.05 ± 0.31 ng/μg [n = 20] vs. 1.43 ± 0.88 ng/μg [n = 20]; P = 0.083). OPN was present in the trabecular meshwork, corneal epithelium and endothelium, iris, ciliary body, retina, vitreous humor, and optic nerve. LFA increased OPN gene expression, but minimal change in OPN protein expression was observed. No difference in intraocular pressure (17.5 ± 2.0 mm Hg [n = 56] vs. 17.3 ± 1.9 mm Hg [n = 68]) but thinner central corneal thickness (91.7 ± 3.6 μm [n = 50] vs. 99.2 ± 5.5 μm [n = 70]) was noted between OPN+/− and wild-type mice.

CONCLUSIONS. OPN is widely distributed in the human eye and was found in lower concentrations in POAG AH. Reduction of OPN in young mice does not affect IOP. (Invest Ophthalmol Vis Sci. 2011;52:6443–6451) DOI:10.1167/iovs.11-7409

Elevated intraocular pressure (IOP) in primary open-angle glaucoma (POAG) is caused by increased outflow resistance within the trabecular meshwork, the tissue that drains most of the aqueous humor (AH) from the anterior segment of the human eye.1 Changes that occur within the trabecular cells, in protein secretions, or within the extracellular matrix have been identified as the most likely causes for increased outflow resistance in POAG.2,3 Traditionally, standard treatment for POAG has focused on reducing IOP surgically, through medications, or both. Pharmaceutical agents such as the prostaglandin analogs (e.g., latanoprost) enhance the flow of AH by stimulating extracellular matrix (ECM) turnover by expression changes of matrix metalloproteinases, tissue inhibitor of metalloproteinases, and matricellular proteins.4–7

Matricellular proteins are nonstructural ECM proteins that regulate cellular function and matrix production by interacting with cellular receptors, growth factors, and cytokines. Matricellular proteins are involved in a wide range of cell modulatory functions, including cell adhesion, migration, survival, and apoptosis.8–11 Several members of the matricellular protein family have been identified as potential regulators of aqueous outflow resistance. For example, mice lacking the matricellular protein SPARC (secreted protein, acidic and rich in cysteine) have lower IOP than wild-type controls.12 Hevin, a matricellular protein with homology to SPARC, interacts with a mutant form of the glaucoma-associated protein myocilin.13 Thrombospondin 1, a multifunctional matricellular protein, is elevated at the mRNA level in primary cultures of lamina cribrosa cells obtained from POAG patients and is also a key activator of transforming growth factor beta (TGFβ), which is elevated in 50% of POAG AH.14–17 Tenascin C, another member of the matricellular protein family, is upregulated in response to mechanical stretch and has been shown to alter cell shape, migration, and adhesion.18–22 However, though somewhat defined, the role of matricellular proteins in IOP regulation and glaucoma is still relatively unknown.

Osteopontin (OPN), another member of the matricellular protein family, is a key regulator of ECM mineralization and bone formation, is considered a strong marker of calcification, vascular diseases (e.g., atherosclerosis), and bone mineralization, and is a prognostic marker for inflammatory heart diseases.23–26 OPN is associated with a wide range of biological processes, including cell adhesion, migration, cell survival, activation of immune cells, and ECM remodeling, and is an inhibitor of vascular calcification.8,9,11,27–29 In the eye, OPN modulates wound healing and neovascularization in the mouse cornea, controls epithelial-mesenchymal transition in the mouse lens, and is upregulated in rat retinal ganglion cells in response to excitotoxic and ischemic injury.30–33 In the normal trabecular meshwork, the expression of OPN mRNA and several other genes associated with calcification inhibition indicates a possible role of these molecules in maintaining a healthy trabecular meshwork.34–36 Imbalance of these inhibitors may lead to a less pliable trabecular meshwork that even-

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tually may affect outflow and IOP. This is consistent with increased trabecular meshwork stiffness in glaucoma.49

We recently identified OPN as one of the most abundant proteins in human AH.40 Given the high level of OPN expression in AH, its potential role as an inhibitor of tissue calcification, and the involvement of other matricellular proteins in IOP regulation, we evaluated the distribution profile of OPN in human ocular tissues and compared OPN concentrations between normal and POAG AH samples. In addition, OPN expression was analyzed in normal human trabecular meshwork (NTM) cells after latanoprost-free acid (LFA) treatment to determine whether OPN levels were influenced by IOP-lowering medications. Finally, the role of OPN in IOP regulation was examined in OPN−/− mice.

**Materials and Methods**

**Collection of Eye Tissues and Fluids**

The protocol for collection of human AH was approved by the Mayo Clinic Institutional Review Board and conforms to tenets of the Declaration of Helsinki.

**Donor.** POAG and normal human donor eyes were obtained from the Minnesota Lions Eye Bank within 12 hours of death. AH was obtained from the anterior chamber with a tuberculin syringe placed parallel to the cornea and was inserted above the limbus. The trabecular meshwork, retina, iris, and optic nerve were dissected and stored at −80°C. Vitreous humor was isolated and centrifuged at 3000g for 10 minutes, and the supernatant was collected and stored at −80°C. Total protein concentrations of all AH, vitreous humor, and tissue lysates were determined using the Bradford protein assay (Bio-Rad, Hercules, CA).

**Surgical.** At the time of trabeculectomy (POAG) or elective cataract surgery, a 30-gauge needle was inserted into the mid-anterior chamber through a paracentesis tract. AH was slowly aspirated until the anterior chamber began to shallow. AH was transferred to a vial and immediately placed in liquid nitrogen. Control samples were obtained from patients with no identified eye disorders other than NTM surgery, a 30-gauge needle was inserted into the mid-anterior chamber through a paracentesis tract. AH was slowly aspirated until the anterior chamber began to shallow. AH was transferred to a vial and immediately placed in liquid nitrogen. Control samples were obtained from patients with no identified eye disorders other than NTM cells after latanoprost-free acid (LFA) treatment to determine whether OPN levels were influenced by IOP-lowering medications. Finally, the role of OPN in IOP regulation was examined in OPN−/− mice.

**Quantification of OPN in AH and Conditioned Media**

OPN was quantified in donor POAG (n = 8), normal donor (n = 9), surgical POAG (n = 20), and elective cataract surgery (n = 20) using a human OPN enzyme-linked immunosorbent assay (ELISA) kit (Quantikine; R&D Systems, Minneapolis, MN; detection limits ≥6 pg/mL). Two microliters of each AH sample was diluted 50-fold with calibrator diluent, mixed with 100 µL assay diluent, and incubated for 2 hours at room temperature in individual microtiter plate wells coated with OPN monoclonal antibody. Each sample was washed three times with 400 µL wash buffer and incubated with an enzyme-linked polyclonal antibody for 2 hours at room temperature. Substrate solution was added to the bound antigen-antibody-enzyme complex, and fluorescence was calculated at 450 nm (with 540 nm as reference wavelength) on a microplate reader (Infinite M200; Tcenan Systems, Inc., San Jose, CA). Concentrations of OPN in POAG and normal samples were established from a standard curve generated using several known concentrations of recombinant human OPN.

For OPN quantification, conditioned media containing 10% FBS was concentrated 10-fold using ultrafiltration spin columns (Vivaspin 6; Sartorius Stedim Biotech GmbH, Goettingen, Germany). Concentrated conditioned media (50 µL) were used for OPN quantification using the human OPN ELISA kit (Quantikine; R&D Systems) as described.

**Western Blot Analysis of Human Eye Tissues and AH**

Various eye tissues dissected from donor eyes were placed in lysis buffer (50 mM Tris, pH 8.0, 0.5% sodium dodecyl sulfate, 0.5% Triton X-100, 137 mM NaCl, 5 mM KCl, 8 mM Na2HPO4·7H2O, 1 mM KH2PO4, protease inhibitors [Roche, Indianapolis, IN]) and homogenized using a rotor stator homogenizer (Polytron; Kinematica, Lucerne, Switzerland). Tissue lysates (15 µg) or AH volume containing 15 µg total protein were mixed in equal proportions with 2× Laemmli buffer, boiled, and separated on 4% to 15% SDS-PAGE gradient gels (Bio-Rad). Proteins were transferred to polyvinylidene difluoride membrane (Milipore, Billerica, MA) in 1× transfer buffer (50 mM Tris, 384 mM glycine, 0.01% SDS, 20% methanol). Membranes were blocked in 20 mM Tris (pH 7.5), 150 mM NaCl, 0.05% Tween-20, and 2% nonfat evaporated milk. Blots were probed with rabbit polyclonal anti-human OPN (Sigma-Aldrich, St. Louis, MO), mouse monoclonal anti-human GAPDH (Novus Biologicals, Littleton, CO), or mouse monoclonal anti-human serum albumin (Abcam, Cambridge, MA) antibodies, followed by a secondary horseradish peroxidase-linked anti-rabbit or anti-mouse antibody (GE Healthcare, Piscataway, NJ). Antibody/antigen complexes were detected using ECL Western blot signal detection reagent (GE Healthcare). Kodak film was used to visualize the protein signals (BioMax XAR; Eastman Kodak, Rochester, NY). Each film was digitized with a photographic scanner (Perfection 2400; Epson, Long Beach, CA). The band intensities in Western blot analysis were normalized to GAPDH (cell lysates) or human serum albumin (in AH) and quantified using ImageJ software (developed by Wayne Rasband, National Institutes of Health, Bethesda, MD; available at http://rsb.info.nih.gov/ij/index.html).

**Immunohistochemistry and Confocal Microscopy**

A human donor eye (81-year-old man) was immersion fixed in 10% neutral buffered formalin within 15 hours of death, dehydrated in a graded series of ethanol (75%, 85%, 95%, and 100%), and embedded in paraffin. Sections (5 µm) were mounted on glass slides (Superfrost/Plus; Fisher, Pittsburgh, PA) and baked at 60°C for 2 hours. For immunohistochemistry, tissue sections were deparaffinized in xylene and rehydrated in a graded series of ethanol (100%, 95%, 80%, 70%) followed by incubation in phosphate-buffered saline. Antigens were retrieved by incubating sections in 1 mM EDTA (pH 8.0) at 95°C for 30 minutes.41 Tissue sections were blocked in phosphate-buffered saline containing 3% bovine serum albumin and 0.1% Triton X-100 and probed with mouse anti-human OPN monoclonal antibody (R&D Systems). Alexa Fluor 488–conjugated goat anti-mouse immunoglobulin was used as a secondary antibody (Molecular Probes, Eugene, OR). Sections were mounted in mounting medium (Vectorshield/DAPI; Vector Laboratories, Burlingame, CA) and examined on a confocal laser microscope (Zeiss 510; Carl Zeiss, Thornwood, NY). Image capture of eye tissues and corresponding controls was performed using the same digital settings.

**Enzymatic Deglycosylation of AH**

An enzymatic protein deglycosylation kit (Sigma-Aldrich) was used for complete removal of O- and N-linked glycosides from AH proteins in accordance with the manufacturer’s instructions. Briefly, 25 µg AH total protein was denatured at 100°C for 5 minutes in the presence of 1× reaction buffer and 2.5 µL denaturation solution in a final volume of 50 µL. The reaction mix was cooled to room temperature, and a panel of five different glycosidases (PNGase F, O-glycosidase, α(2,3,6,8,9)neuraminidase, β(1−3)galactosidase and β-N-acetylglucosaminidase) was added with 2.5 µL nonionic surfactant (Triton X-100; Sigma-Aldrich). The reaction mix was incubated for 3 hours at 37°C. After incubation, 35 µL reaction mixture was combined with 5× Laemmli buffer, boiled, and separated on 4% to 15% SDS-PAGE gradient gels. Proteins were transferred to polyvinylidene difluoride membrane and probed with OPN antibody, as described.
Primary Culture of Human Trabecular Meshwork Cells

Three primary human NTM cell lines derived from independent donor eyes (20- and 46-year-old women; 1-year-old boy) were isolated as described previously. Briefly, trabecular meshwork tissue was dissected from human donor eyes and digested with collagenase A (Worthington Biochemical Corporation, Lakewood, NJ). The cells were pelleted, resuspended in Dulbecco’s modified Eagle’s medium (DMEM; Mediatech, Inc., Herndon, VA), and supplemented with 10% fetal bovine serum (FBS; Cellgro, Manassas, VA) and penicillin-streptomycin (100 U/ml; final concentration; Invitrogen, Carlsbad, CA). Cells were placed in a six-well plate and were grown to confluence at 37°C and 5% CO2. Cells from passages 3 to 7 were used for experiments performed in this study.

LFA Treatment of NTM Cell Cultures

LFA was purchased as a solution in methyl acetate from Cayman Chemical (Ann Arbor, MI). For use in the experiments, methyl acetate was evaporated under a gentle stream of nitrogen, and LFA was reconstituted in ethanol and purged with nitrogen at a concentration of 10−7 M. The study solution of LFA was diluted in DMEM with or without 10% FBS to a final working concentration of 10−7 M.

Primary human NTM cell lines (n = 3) were grown to confluence in six-well plates containing DMEM supplemented with 10% FBS. Confluent cells were washed twice with PBS and incubated in serum-free DMEM for 24 hours to synchronize the growth potential of the cells. After overnight serum deprivation, cells were treated with LFA (10−7 M final concentration) and vehicle control (ethanol, final dilution 1:1000) in DMEM with and without 10% FBS. Cells and conditioned media were harvested 24, 48, and 72 hours after treatment.

Total RNA. Total RNA was isolated using a purification kit (RNasy Mini Kit; Qiagen, Valencia, CA) according to the manufacturer’s protocol. Total RNA (250 ng) was reverse transcribed into cDNA using a cDNA synthesis kit (Script; Bio-Rad). Quantitative real-time polymerase chain reaction using OPN (forward, 5′-ACAGCCAGGGACTCCATG-3′; reverse, 5′-ACACTATACCCTGCGATC-3′) and GAPDH (forward, 5′-CCTCTGAATTCAAGC-3′; reverse, 5′-GGTAGCAAAATGTTGT-3′) primers was performed (LightCycler 480 SYBR Green Master Kit; Roche) by pre-denaturation at 95°C (1 cycle) followed by amplification (45 cycles): denaturation at 95°C; annealing at 63°C; and extension at 72°C. Fold change was calculated after normalization with GAPDH.

Total Protein. NTM cell pellets were suspended in ice-cold lysis buffer and passed repeatedly through a 21-gauge needle. Lysate was centrifuged at 13,000g for 10 minutes, and total protein was quantified by the Bradford assay. Equal protein amounts (15 μg) were separated on 4% to 15% SDS-PAGE gels, transferred to polyvinylidene difluoride membrane, and probed with OPN antibody as described.

Measurement of IOP in OPN+/−/− Mice

Adult OPN+/− mice (C57BL/6) and wild-type controls were purchased from the Jackson Laboratory (Bar Harbor, ME). The mice were anesthetized by intraperitoneal injection of a ketamine/xylazine mixture (100 and 9 mg/kg, respectively; Phoenix Pharmaceutical, St. Joseph, MO). A previously validated commercial rebound tonometer (TonoLab; Colonial Medical Supply, Franconia, NH) was used to take three sets of six IOP measurements from each eye. Right and left eyes were measured alternately, with the initial eye selected randomly. All measurements were taken between 4 and 7 minutes after intraperitoneal injection because previous studies have shown this to be a period with stable IOP. As indicated by the manufacturer, the tonometer (TonoLab; Colonial Medical Supply) was fixed horizontally for all measurements, and the tip of the probe was placed 2 to 3 mm from the eye. To reduce variability in measurements, the tonometer was modified to include a pedal that activated the probe without the need to handle the device. The probe contacted the eye perpendicularly over the central cornea. Verification of targeting was performed using direct visualization using 5.5× magnification. The average of a set of measurements for each eye was accepted only if the device indicated that there was “no significant variability” (per the protocol manual; Colonial Medical Supply). Measurements were taken at the age of 7 weeks between 11:00 and 15:00 hours to minimize the effect of diurnal variation.

Measurement of Central Corneal Thickness

In humans, contact tonometry measurements by variable force spring tonometers (e.g., Goldmann, Tono-Pen) may be affected by central corneal thickness. To assess whether this confounder contributes to any observed IOP difference, the central corneal thickness of 6.5-week-old OPN+/−/− and wild-type mice were measured using optical coherence tomography (Stratus; Carl Zeiss Meditec, Inc., Dublin, CA). The mice were imaged with optical coherence tomography under anesthesia (ketamine/xylazine mixture). Central corneal thickness was obtained by using the internal software (version 4.0.7) and measuring the distance between the epithelial and endothelial peak amplitudes.

Histology

Eyes were enucleated from euthanized 8-week-old wild-type and OPN+/−/− mice and were placed in 2.5% glutaraldehyde/2% formaldehyde in 0.1 M cacodylate buffer containing 0.08 M CaCl2 at 4°C for 20 minutes. After incubation, eyes were hemisected, and the tissue minus the lens was returned to fixative for 24 hours. Tissue was postfixed for 1.5 hours in 2% aqueous OsO4. Tissue was dehydrated in graded concentrations of ethanol, transitioned in propylene oxide, infiltrated with propylene oxide and Epon mixtures (TAAB 812 resin; Marivac, Quebec, Canada), embedded in Epon, and cured for 48 hours at 60°C. Sections (1 μm) were stained with 1% toluidine blue in 1% borate buffer.

Statistical Analysis

Results are expressed as mean ± SD. Data from various experimental and control groups were compared using Student’s unpaired t-test. Differences were considered significant when P < 0.05.

RESULTS

Expression of OPN in Normal and POAG Human AH

Previous proteomic studies from our laboratory have shown high levels of OPN in human AH. To validate this, OPN was evaluated in donor POAG and normal AH by ELISA. In AH obtained from donor eyes, POAG OPN levels (normalized to total protein concentration) were significantly lower than normal (0.54 ± 0.18 ng/μg vs. 0.77 ± 0.23 ng/μg; P = 0.039; Fig. 1A). No statistical difference between total protein of donor POAG and normal AH was observed (0.929 ± 0.28 μg/μL vs. 0.919 ± 0.29 μg/μL; P = 0.96). When analyzed by volume, OPN concentration was also found to be significantly lower in donor POAG AH compared with normal AH (0.5 ± 0.18 ng/μL vs. 0.67 ± 0.14 ng/μL; P = 0.044).

OPN levels in AH obtained during surgery showed a similar trend to OPN levels in donor eyes. In surgical POAG AH, OPN levels (normalized to total protein concentration) were lower than in AH from elective cataract controls (1.05 ng/μL vs. 1.43 ng/μg; P = 0.083; Fig. 1B). No significant change in OPN concentration by volume was noted between surgical POAG and elective cataract control AH samples (0.27 ± 0.09 μg/mL vs. 0.26 ± 0.16 μg/μL; P = 0.80). However, total protein concentration was significantly higher in surgical POAG AH than in elective cataract control AH (0.255 ± 0.08 μg/μL vs. 0.195 ± 0.09 μg/μL; P = 0.005).

To confirm the ELISA results, Western blot analysis was performed on POAG and normal AH obtained from donor eyes. Using
an anti–OPN polyclonal antibody, six OPN protein products were detected (Fig. 2A). Results show a wide variation in OPN expression profiles within POAG and normal samples. Interestingly, OPN band 4 (70 kDa) was found to be absent in 3 of the 6 POAG samples (Fig. 2A, lanes 10, 12, 13). Although band 4 was present in all normal samples, reduced expression was noted in 2 of the 8 samples (Fig. 2A, lanes 4 and 5). The mean intensity of individual bands 1 to 5 (normalized to HAS) was found to be lower in POAG than in normal (Fig. 2B) AH. Conversely, band 6 was slightly higher in POAG than in normal AH. Only band 2 was found to be significantly different between POAG and normal AH (P < 0.02). When the intensity of all six bands was combined, POAG samples had lower OPN expression than normal samples, consistent with ELISA results (Fig. 1).

**Evaluation of OPN Expression in Ocular Tissues by Immunohistochemistry**

Because several components of AH are actively secreted by eye tissues, there is a high probability that OPN is synthesized by eye tissues themselves. To analyze this, we evaluated the presence of OPN by immunohistochemistry in various ocular tissues. Using a monoclonal anti–OPN antibody, OPN staining was shown to be present in the trabecular meshwork, corneal epithelium and endothelium, iris, ciliary body, retina and anterior and posterior optic nerves (Fig. 3).

**OPN Expression Pattern in Various Ocular Tissues**

The expression profile of OPN was further evaluated by Western blot analysis in iris, retina, optic nerve, vitreous humor, and trabecular meshwork obtained from three normal donor eyes (Fig. 4). Although there was variation in OPN expression among the samples, it was found to be strongly expressed in retina, trabecular meshwork, and optic nerve, confirming expression studies performed by immunohistochemistry. Only 1 of 3 samples showed OPN in the iris, and 2 of 3 samples showed OPN in vitreous humor.

OPN protein size also varied among tissues (Fig. 4). In retina and vitreous humor, a doublet was observed at approximately 55 kDa. In the TM, OPN was present as a single band at 55 kDa. OPN expression in optic nerve varied in each sample, appearing as either a singlet or a doublet at 55 kDa. Smaller molecular weight OPN proteins were identified in the retina (35 kDa, 26 kDa, 25 kDa), optic nerve (35 kDa), and vitreous (35 kDa, 26 kDa).
We hypothesized that at least some of the multiple OPN bands were attributed to differential glycosylation of the protein because several investigators have shown extensive glycosylation in OPN. Enzymatic deglycosylation of the protein component in AH from normal and POAG donor eyes showed a downward shift of several OPN bands by Western blot (Fig. 5A). This suggests that several glycosylated products of OPN are present in AH. In contrast, enzymatic deglycosylation of optic nerve and TM lysate showed no change in OPN migration in POAG and normal samples, indicating a lack of glycosylation (Fig. 5B).

**Evaluation of OPN Expression in Response to LFA in Cultured NTM Cells**

One of the limitations of studying proteins in POAG AH is that most of the samples are from patients who are on pressure-lowering medications such as prostaglandin analogs. To deter-
Given that OPN was differentially expressed in POAG AH and SPARC−/− mice had lower IOP than littermate controls, we investigated whether the lack of OPN had any effect on IOP. Analysis of IOP in 7-week-old OPN−/− mice showed no difference in IOP when compared with wild-type controls (17.5 ± 2.0 mm Hg [n = 56] vs. 17.3 ± 1.9 mm Hg [n = 68]). Optical coherence tomography showed OPN−/− mice had thinner central corneal thickness than did wild-type control (91.7 ± 3.6 μm [n = 50] vs. 99.2 ± 5.5 μm [n = 70]; P > 0.001). There were no observable anatomic differences within the outflow tissues between the wild-type and the OPN−/− mice at the light microscopy level (Fig. 7).

**DISCUSSION**

IOP regulation within the human eye occurs primarily in the conventional outflow pathway, where cellular components of the trabecular meshwork and Schlemm’s canal work in conjunction with the ECM to regulate AH removal from the anterior segment. Associated with the interplay between the cells and the extracellular matrix are matricellular proteins. SPARC, a member of the matricellular protein family, has been shown to be directly involved in IOP regulation.12 Conversely, hevin, a close homolog of SPARC, does not have a direct effect on IOP.55 In this study, we found the matricellular protein OPN to be expressed at lower levels in POAG AH than in controls. Variation in OPN protein size was observed in POAG AH. Some, but not all, of the OPN size variation was due to glycosylation. However, unlike SPARC−/− mice, we did not observe any significant difference in IOP of OPN−/− mice when compared with wild-type controls.

Our study indicates a decreasing trend of OPN in POAG AH, whereas other matricellular proteins are reported to be upregulated at the mRNA level in POAG after mechanical stretch or in response to elevated IOP.22,53,54 The reason for decreased OPN levels in POAG AH is unknown. Previous studies evaluating gene expression of human TM cells in response to glaucomatous insults such as dexamethasone treatment did not find upregulation of the OPN gene.55,56 However, contrasting reports from nonocular cells (osteoprecursor and lung cells) suggest that dexamethasone treatment may affect OPN gene expression in a cell type–specific manner.57,58 Unlike other matricellular proteins, OPN is an inhibitor of calcification and aids in the dissolution of mineral deposits.59,60 Matrix Gla protein, another inhibitor of calcification, is decreased in the TMs isolated from POAG patients.38 With OPN decreased in POAG AH and matrix Gla protein reduced in POAG TMs, it is
OPN wild-type and (B) age-matched normal corneal angles of mice. (C) Intracellular and (D) secreted OPN protein levels did not change within 48 hours after LFA treatment in 10% FBS-containing media. A slight increase (20%) in secreted OPN was found at 72 hours but was not significant.

tempting to speculate that a decrease in calcification inhibitors may lead to an increase in TM mineralization. Normally, the trabecular meshwork is a spongiform tissue that reacts to various forms of stress (e.g., shear, stretch) to modulate outflow. An increase in mineralization may result in a TM that is stiffer and less responsive to stress, leading to decreased outflow and elevated IOP. Last et al.\(^5\) reported that the TM is less pliable and stiffer in POAG than in age-matched normal controls, supporting this concept.

Another possibility for lower OPN levels in POAG AH may be the presence of a negative feedback loop by which the eye responds to increased IOP.\(^6\) Matrix metalloproteinases have often been shown to be expressed in an opposing manner in physiological or pathologic processes. In the case of non-small cell lung cancer, increased OPN favors progression of the disease but was accompanied by a lower expression of SPARC.\(^6\) In addition, overexpression of OPN mRNA has been linked to a poor prognosis for ovarian cancer patients, whereas overexpression of SPARC protein in an ovarian carcinoma cell line is inversely correlated with the degree of malignancy by promoting apoptosis, indicating a tumor suppressor role for SPARC in these cells.\(^6\)

Contradictory to our results is the finding of increased OPN in the AH of DBA2/J mice.\(^5\) Although this may be explained by the differences between humans and mice, it may also represent a difference in the etiology of the glaucoma. In DBA2/J mice, elevated IOP is caused by the shedding of pigment from the iris, resulting in a significant inflammatory response and eventual neovascular scarring of the angle. In POAG, pigment is generally not a factor, but the etiology of elevated IOP is unknown.

Our finding of upregulated OPN mRNA in response to LFA treatment corroborates a previous report showing prostaglandin treatment increased OPN gene expression in human adipose tissue-derived mesenchymal stem cells.\(^6\) In our study, we observed an increase in OPN mRNA after LFA treatment. However, we did not observe a change in OPN protein expression within 48 hours of LFA treatment (cellular or secreted). Only at 72 hours after treatment did we see a slight increase in OPN secretion. This suggests that LFA does not contribute to the lower OPN concentrations seen in POAG AH. Interestingly, we could induce an increase in OPN gene transcription only in NTM cells treated with LFA in serum-containing media. Whether LFA needed auxiliary proteins from the FBS to induce OPN transcription is unknown.

In humans, OPN is found in many tissues and almost all body fluids, including blood, urine, and milk.\(^5\) Extensive post-translational modifications affect OPN function in a tissue-specific manner.\(^5\) In AH, six different OPN products were identified; sizes ranged from 55 to 180 kDa. OPN is glycosylated in AH, but these modifications do not account for all the OPN size variations. In addition to glycosylation, OPN is post-translationally modified by sulfation, serine/threonine phosphorylation, sialylation, and transglutamination.\(^5\) The higher molecular weight bands (\(>55\) kDa) we observed in AH corroborate with similarly sized OPN fragments found in bone lysate and calcified arteries of mice.\(^6\) Bands 4 and 5 correspond to full-length OPN (Fig. 2). Reduction of band 4 in POAG AH suggests a loss of OPN full-length biological function. The remaining bands are smaller than full-length OPN, consistent with cleaved products. OPN has been shown to be cleaved by a variety of molecules, including thrombin, plasmin, cathepsin D, and various matrix metalloproteinases.\(^5\) The importance of the smaller fragments is unknown. Given the inadequate understanding of the pathophysiology of glaucoma, it is difficult to say whether changes in OPN are the cause or the effect of the disease itself. Further experiments are required to
determine the biological function of OPN in normal AH and the consequence of reduction in several OPN fragments.

In eye tissues, differences in OPN banding patterns were also noted in trabecular meshwork, iris, optic nerve, retina, vitreous humor, and AH. In trabecular meshwork, OPN was seen as a single band found at 55 kDa. Although the 55-kDa band was consistent between all the tissues analyzed, a doublet was found in the retina, optic nerve, and vitreous humor in at least 1 of the 3 samples studied. The observed differences in OPN expression in different eye tissues may indicate active synthesis of the protein and may contribute to the total secreted OPN in AH. However, the contribution of OPN via serum (at least for some of the OPN forms or part of the total OPN) cannot be ruled out.

Wild-type mice have a wide range of baseline IOP (4–20 mm Hg) that is strain dependent.12,44,63,73,74 Thus, it is important that comparative studies, such as ours, use paired wild-type and knockout mice from the same background strain. Unlike SPARC−/− mice, which had lower IOP than wild-type controls,12 the OPN−/− mice did not show any change in IOP compared with controls. OPN−/− mice did, however, have a 7.6% thinner central corneal thickness. In humans, a 10% change in central corneal thickness is generally considered significant using Goldmann applanation tonometry. However, our calibration with manometry indicates that central corneal thickness does not appear to have a significant impact with rebound tonometry.48 Previous studies in mice found a high correlation between the IOPs measured by tonometer and manometry. This suggests that central corneal thickness does not affect IOP measurements using rebound tonometry.63,74 Taken together, these results indicate that there may not be a direct involvement of OPN in IOP regulation in young mice.

In summary, our results demonstrate a strong presence of OPN in various tissues of the eye and an overall decrease of OPN in POAG AH compared with control AH. Further studies are required to specifically determine whether OPN can be used as a potential marker for POAG.

References

Osteopontin in POAG


65. Christensen B, Nielsen MS, Haselmann KF, Petersen TE, Sorenson ES. Osteopontin is cleaved at multiple sites close to its integrin-binding motifs in milk and is a novel substrate for plasmin and cathepsin D. J Biol Chem. 2010;285:7929–7937.


