Extracellular Matrix Stiffness and TGFβ2 Regulate YAP/TAZ Activity in Human Trabecular Meshwork Cells

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Primary open-angle glaucoma progression is associated with increased human trabecular meshwork (HTM) stiffness and elevated transforming growth factor beta 2 (TGFβ2) levels in the aqueous humor. Increased transcriptional activity of Yes-associated protein (YAP) and transcriptional coactivator with PDZ-binding motif (TAZ), central players in mechanotransduction, are implicated in glaucomatous HTM cell dysfunction. Yet, the detailed mechanisms underlying YAP/TAZ modulation in HTM cells in response to alterations in extracellular matrix (ECM) stiffness and TGFβ2 levels are not well understood. Using biomimetic ECM hydrogels with tunable stiffness, here we show that increased ECM stiffness elevates YAP/TAZ nuclear localization potentially through modulating focal adhesions and cytoskeletal rearrangement. Furthermore, TGFβ2 increased nuclear YAP/TAZ in both normal and glaucomatous HTM cells, which was prevented by inhibiting extracellular-signal-regulated kinase and Rho-associated kinase signaling pathways. Filamentous (F)-actin depolymerization reversed TGFβ2-induced YAP/TAZ nuclear localization. YAP/TAZ depletion using siRNA or verteporfin decreased focal adhesions, ECM remodeling and cell contractile properties. Similarly, YAP/TAZ inactivation with verteporfin partially blocked TGFβ2-induced hydrogel contraction and stiffening. Collectively, our data provide evidence for a pathologic role of aberrant YAP/TAZ signaling in glaucomatous HTM cell dysfunction, and may help inform strategies for the development of novel multifactorial approaches to prevent progressive ocular hypertension in glaucoma.

Keywords: mechanotransduction, hydrogel, HTM cell contractility, HTM stiffness, POAG
INTRODUCTION

Primary open-angle glaucoma (POAG) is a leading cause of irreversible vision loss worldwide, and elevated intraocular pressure (IOP) is the primary modifiable risk factor (Quigley, 1993; Quigley and Broman, 2006; Kwon et al., 2009; Tham et al., 2014; Tamm et al., 2015). Elevated IOP results from increased resistance to aqueous humor outflow in the juxtacanalicular region of the conventional outflow pathway where the trabecular meshwork (TM) and Schlemm’s canal inner wall cells interact (Brubaker, 1991). Human trabecular meshwork (HTM) cells within the juxtacanalicular tissue are surrounded by a complex extracellular matrix (ECM), which is primarily composed of non-fibrillar and fibrillar collagens, elastic fibrils, proteoglycans and the glycosaminoglycan hyaluronic acid (Acott and Kelley, 2008; Tamm, 2009; Hann and Fautsch, 2011; Keller and Acott, 2013; Abu-Hassan et al., 2014). In addition to providing structural support, the ECM imparts biochemical signals (e.g., hormones, growth factors and diffusible morphogens) and mechanical cues (e.g., matrix stiffness; tensile, compressive and shear forces; topographical strain) to regulate cell behaviors (Frantz et al., 2010). These external biophysical cues are translated into internal biochemical signaling cascades through a process known as mechanotransduction, and cells play an active role in remodeling their matrix to promote mechanical homeostasis and maintain tissue-level functionality (Lampi and Reinhart-King, 2018).

HTM cells are exposed to a variety of biophysical cues through cell-cell and cell-ECM interactions, and fluctuations in IOP. In POAG, impaired HTM cell function (i.e., remodeling of cell cytoskeleton, increased cell stiffness) and increased ECM deposition contribute to HTM stiffening (Schlunck et al., 2008; Han et al., 2011; McKee et al., 2011; Wang et al., 2017b; Li et al., 2021a). Tissue stiffening affects HTM cell function and IOP in a feed-forward cycle characterized by dynamic reciprocity. HTM stiffening in POAG indicates that a biophysical component likely contributes to IOP regulation. Therefore, targeting mechanotransduction pathways in the HTM to interrupt this feed-forward cycle between HTM stiffening and cellular response to the stiffened ECM is emerging as a promising strategy for managing ocular hypertension.

Yes-associated protein (YAP) and transcriptional coactivator with PDZ-binding motif (TAZ, encoded by WWTR1) are central players in mechanotransduction. YAP/TAZ can be activated by PDZ-binding motif (TAZ, encoded by WWTR1) are central players in mechanotransduction. YAP/TAZ can be activated by increased ECM stiffness, mechanical stress and growth factors (Dupont et al., 2011; Boopathy and Hong, 2019). It has been demonstrated that certain glaucoma related molecules that influence outflow function, such as dexamethasone, lysophosphatidic acid and interleukin-6, mediate the expression of YAP and TAZ in HTM cells (Ho et al., 2018; Honjo et al., 2018; Peng et al., 2018; Yemanyi and Raghunathan, 2020). YAP was found to be reduced in HTM cells incubated on stiff two-dimensional (2D) polyacrylamide hydrogels, while TAZ levels were increased (Raghunathan et al., 2013). Other research showed that both YAP and TAZ were significantly higher in HTM cells on stiffer polyacrylamide substrates or stiffened HTM cell-derived matrices (Thomasy et al., 2013; Yemanyi et al., 2020).

Whilst it appears that these findings may be contradictory, it is important to note that functional outcomes of YAP and TAZ are governed by their subcellular localization which is substrate and context dependent. Together, these highlight the importance of studying YAP/TAZ signaling in a microenvironment mimicking the native tissue to further our understanding of HTM cell mechanobiology with an emphasis on cell-ECM interactions. This is especially important considering a recent multi-ethnic genome wide meta-analysis report that identified YAP1 as a potential genetic risk factor for POAG across European, Asian, and African ancestries implicating a causal relationship for outflow dysfunction (Gharahkhani et al., 2021).

Biomaterials for cell culture applications are designed to mimic aspects of the ECM and provide researchers with methods to better understand cell behaviors. Protein-based hydrogels (i.e., water-swollen networks of polymers) are attractive biomaterials owing to their similarity to in vivo cellular microenvironments, biocompatibility, biodegradability and tunable mechanical properties (Drury and Mooney, 2003; Panahi and Baghban-Salehi, 2019). Recently, we developed a bioengineered hydrogel composed of HTM cells and ECM biopolymers found in the native HTM tissue, and demonstrated its viability for investigating cell-ECM interactions under normal and simulated glaucomatous conditions in both 2D and 3D conformations (Li et al., 2021a; Li et al., 2021b).

Here, we hypothesized that altered YAP/TAZ activity drives HTM cell dysfunction under simulated glaucomatous conditions. The TM from POAG eyes is stiffer than that from healthy eyes (Last et al., 2011; Wang et al., 2017a; Vahabikashi et al., 2019); transforming growth factor beta 2 (TGFβ2), the predominant TGFβ isoform in the eye and aqueous humor, has been identified as a major contributor to the pathologic changes occurring in ocular hypertension and POAG (Granstein et al., 1990; Quigley, 1993; Inatani et al., 2001; Fuchshofer and Tamm, 2009; Agarwal et al., 2015; Kasetti et al., 2018). Using TGFβ2 as a glaucomatous stimulus and tuning the stiffness of our hydrogels, here, we investigated the effects of ECM stiffening and TGFβ2 on regulating YAP/TAZ in HTM cells in 2D and 3D cultures. In this tissue-mimetic environment, we then investigated whether YAP/TAZ inhibition would alleviate ECM stiffness- or TGFβ2-induced HTM cell pathobiology.

MATERIALS AND METHODS

HTM Cell Isolation and Culture

Experiments using human donor eye tissue were approved by the SUNY Upstate Medical University Institutional Review Board (protocol #1211036), and were performed in accordance with the tenets of the Declaration of Helsinki for the use of human tissue. Primary HTM cells were isolated from healthy donor corneal rims discarded after transplant surgery as recently described (Li et al., 2021a; Li et al., 2021b), and cultured according to established protocols (Stamer et al., 1995; Keller et al., 2018). Five HTM cell strains (HTM05, HTM12, HTM14, HTM17, HTM19) were used for the experiments in this study.
(Supplementary Table S1). All HTM cell strains were validated with dexamethasone (DEX; Fisher Scientific, Waltham, MA, United States; 100 nM) induced myocilin expression in more than 50% of cells by immunocytochemistry and immunoblot analyses [Supplementary Figures S1A,B and (Li et al., 2021a; Li et al., 2021b)]. Different combinations of two to three HTM cell strains were used per experiment with three to four replicates each, depending on cell availability, and all studies were conducted between cell passage 3–7. HTM cells were cultured in low-glucose Dulbecco’s Modified Eagle’s Medium (DMEM; Gibco; Thermo Fisher Scientific) containing 10% fetal bovine serum (FBS; Atlanta Biologicals, Flowery Branch, GA, United States) and 1% penicillin/streptomycin/glutamine (P(SG; Gibco), and maintained at 37°C in a humidified atmosphere with 5% CO₂, Fresh media was supplied every 2–3 days.

Glaucomatic HTM Donor History
Glaucomatic HTM cells (GT M211 and GT M1445) were used in this study. GTM211 was isolated from a 75-year-old, white female that was taking Xalatan (0.005%) in both eyes nightly for treatment of ocular hypertension. The patient was diagnosed with glaucoma, and had bilateral cataract surgery, and retina surgery. Upon receipt of donor eyes by W.D.S., outflow facility measurements during constant pressure perfusion conditions were 0.24 and 0.17 μL/min/mmHg from OD and OS eyes, respectively. GTM211 were isolated from the OS eye of the donor and characterized as previously described (Li et al., 2021a). GTM1445 has been characterized in our previous study (Li et al., 2021a). GTM cell strain information can be found in Supplementary Table S1.

Preparation of Hydrogels
Hydrogel precursors methacrylate-conjugated bovine collagen type I [MA-COL, Advanced BioMatrix, Carlsbad, CA, United States; 36 mg/ml (all final concentrations)], thiol-conjugated hyaluronic acid (SH-HA, Glycosil®, Advanced BioMatrix; 0.5 mg/ml, 0.025% (w/v) photoinitiator Irgacure® 2959; Sigma-Aldrich, St. Louis, MO, United States) and in-house expressed elastin-like polypeptide (ELP, thiol via KCTS flanks (Li et al., 2021a); 2.5 mg/ml) were thoroughly mixed. Thirty microliters of the hydrogel solution were pipetted onto a Surfasil (Fisher Scientific) coated 12-mm round glass coverslip followed by placing a regular 12-mm round glass coverslip onto the hydrogels. Constructs were crosslinked by exposure to UV light (OmniCure S1500 UV Spot Curing System; Excelitas Technologies, Mississauga, Ontario, Canada) at 320–500 nm, 2.2 W/cm² for 5 s, as previously described (Li et al., 2021a). The hydrogel-dermed coverslips were removed with fine-tipped tweezers and placed in 24-well culture plates (Corning; Thermo Fisher Scientific). Hydrogel stiffening was achieved by soaking the hydrogels in 0.1% (w/v) riboflavin (RF; Sigma) for 5 min, followed by low-intensity secondary UV crosslinking (bandpass filter: 405–500 nm, 5.4 mW/cm²) for 5 min.

HTM Cell Treatments
HTM/GTM cells were seeded at 2 × 10⁴ cells/cm² on premade hydrogels/cover slips/tissue culture plates, and cultured in DMEM with 10% FBS and 1% PSG for 1 or 2 days. Then, HTM cells were cultured in serum-free DMEM with 1% PSG and subjected to the different treatments for 3 days: TGFβ2 (2.5 ng/ml; R&D Systems, Minneapolis, MN, United States), the ERK inhibitor U0126 (10 μM; Promega, Madison, WI, United States), the Rho-associated kinase (ROCK) inhibitor Y27632 (10 μM; Sigma-Aldrich), the actin de-polymerizer latrunculin B (10 μM; Tocris Bioscience; Thermo Fisher Scientific), or the YAP inhibitor verteporfin (0.5 μM; Sigma). The monolayer HTM cells were processed for immunoblot, qRT-PCR and immunocytochemistry analyses.

Immunoblot Analysis
Protein was extracted from cells using lysis buffer (CellLytic™ M, Sigma-Aldrich) supplemented with Halt™ protease/phosphatase inhibitor cocktail (Thermo Fisher Scientific). Equal protein amounts (10 μg), determined by standard bicinchoninic acid assay (Pierce; Thermo Fisher Scientific), in 4× loading buffer (Invitrogen; Thermo Fisher Scientific) with 5% beta-mercaptoethanol (Fisher Scientific) were boiled for 5 min and subjected to SDS-PAGE using NuPAGE™ 4–12% Bis-Tris Gels (Invitrogen; Thermo Fisher Scientific) at 120 V for 80 min and transferred to 0.45 μm PVDF membranes (Sigma; Thermo Fisher Scientific). Membranes were blocked with 5% bovine serum albumin (Thermo Fisher Scientific) in tris-buffered saline with 0.2% Tween® 20 (Thermo Fisher Scientific), and probed with various primary antibodies followed by incubation with HRP-conjugated secondary antibodies or fluorescent secondary antibodies (LI-COR, Lincoln, NE, United States). Bound antibodies were visualized with the enhanced chemiluminescent detection system (Pierce) on autoradiography film (Thermo Fisher Scientific) or Odyssey® CLx imager (LI-COR). Densitometry was performed using the open-source National Institutes of Health software platform, FIJ I (Schindelin et al., 2012) or Image Studio™ Lite (LI-COR); data were normalized to GAPDH. A list of all of antibodies used in this study, including their working dilutions, can be found in Supplementary Table S2.

Immunocytochemistry Analysis
HTM cells in presence of the different treatments were fixed with 4% paraformaldehyde (Thermo Fisher Scientific) at room temperature for 20 min, permeabilized with 0.5% Triton™ X-100 (Thermo Fisher Scientific), blocked with blocking buffer (BioGenEx), and incubated with primary antibodies, followed by incubation with fluorescent secondary antibodies; nuclei were counterstained with 4',6'-diamidino-2-phenylindole (DAPI; Abcam). Similarly, cells were stained with Phalloidin-iFluor 488 or 594 (Abcam)/DAPI according to the manufacturer’s instructions. Coverslips were mounted with ProLong™ Gold Antifade (Invitrogen) on Superfrost™ microscope slides (Fisher Scientific), and fluorescent images were acquired with an Eclipse NI microscope (Nikon Instruments, Melville, NY, United States) or a Zeiss LSM 780 confocal microscope (Zeiss, Germany). Fluorescent signal intensity was measured using FIJI software. A list of
all of antibodies used in this study, including their working dilutions, can be found in Supplementary Table S2.

Image Analysis

All image analysis was performed using FIJI software. Briefly, the cytoplasmatic YAP intensity was measured by subtracting the overlapping nuclear (DAPI) intensity from the total YAP intensity. The nuclear YAP intensity was recorded as the proportion of total YAP intensity that overlapped with the nucleus (DAPI). YAP/TAZ nuclear/cytoplasmic (N/C) ratio was calculated as follows: N/C ratio = (nuclear YAP signal/area of nucleus)/(cytoplasmic signal/area of cytoplasm). Fluorescence intensity of F-actin, α-SMA, FN, and p-MLC were measured in at least 20 images from 2 HTM cell strains with three replicates per HTM cell strain with image background subtraction using FIJI software. For number of vinculin puncta quantification, the 3D Object Counter plugin in FIJI software was used to threshold images and quantify puncta/0.05 mm² (above 2 μm in size). Nuclear area was also measured using FIJI software. At least 100 nuclei were analyzed from 2 HTM cell strains with three replicates per HTM cell strain.

Quantitative Reverse Transcription-Polymerase Chain Reaction Analysis

Total RNA was extracted from HTM cells using PureLink RNA Mini Kit (Invitrogen). RNA concentration was determined with a NanoDrop spectrophotometer (Thermo Fisher Scientific). RNA was reverse transcribed using iScript™ cDNA Synthesis Kit (BioRad, Hercules, CA, United States). One hundred nanograms of cDNA were amplified in duplicates in each 40-cycle reaction using a CFX 384 Real Time PCR System (BioRad) with annealing temperature set at 60°C, Power SYBR® Green PCR Master Mix (Thermo Fisher Scientific), and custom-designed qRT-PCR primers. Transcript levels were normalized to GAPDH, and mRNA fold-changes calculated relative to mean values of normal HTM cell controls using the comparative C₅₇ method (Schmittgen and Livak, 2008). A list of all of primers used in this study can be found in Supplementary Table S3.

Active TGFβ2 Enzyme-Linked Immunosorbert Assay

TGFβ2 levels were quantified using the Quantikine ELISA kit (R&D Systems). HTM and GTM cells were seeded at 2 × 10⁴ cells/cm² on tissue culture plates, and cultured in DMEM with 10% FBS and 1% PSG in presence of the different treatments. Longitudinal bright field images were acquired at 0 and 5 days with an Eclipse Ti microscope (Nikon). Construct area from n = 8–11 hydrogels per group from 2 HTM/GTM cell strains with three to four replicates per HTM/GTM cell strain was measured using FIJI software and normalized to 0 days followed by normalization to controls.

siRNA Transfection

HTM cells were depleted of YAP and TAZ using siRNA-loaded lipofectamine RNAiMax (Invitrogen) according to the manufacturer’s instructions. In brief, HTM cells were seeded at 2 × 10⁴ cells/cm² on RF double-crosslinked hydrogels in DMEM with 10% FBS and 1% PSG. The following day, the cell culture medium was changed to antibiotic-free and serum-free DMEM and the samples were kept in culture for 24 h followed by transfection. Transfection was performed using a final concentration 3% (v/v) lipofectamine RNAiMax with 150 nM RNAi duplexes (custom oligonucleotides; Dharmacon). Transfected HTM cells were used 48 h after transfection. ON-TARGET plus nontargeting siRNA were obtained from Dharmacon. Custom siRNA were based on validated sequences previously described (Dupont et al., 2011): YAP, sense, 5′-GACAUCUUCUGGUCAGAGA-3′, and YAP, anti-sense, 5′-UCUCUGACCAAGAAGUG-3′; TAZ, sense, 5′-ACGUUGACUUAGGAACUU-3′, and TAZ, anti-sense, 5′-AAAGUUCUAAGUCAACG-3′.

HTM Hydrogel Contraction Analysis

HTM cell-laden hydrogels were prepared by mixing HTM cells (1.0 × 10⁶ cells/ml) with MA-COL (3.6 mg/ml [all final concentrations]), SH-HA (0.5 mg/ml, 0.025% [w/v] photoinitiator) and ELP (2.5 mg/ml on ice, followed by pipetting 10 µl droplets of the HTM cell-laden hydrogel precursor solution onto polydimethylsiloxane (PDMS; Sylgard 184; Dow Corning) coated 24-well culture plates. Constructs were crosslinked as described above (320–500 nm, 2.2 W/cm², 5 s). HTM cell-laden hydrogels were cultured in DMEM with 10% FBS and 1% PSG in presence of the different treatments. Longitudinal brightfield images were acquired at 0 and 5 days with an Eclipse Ti microscope (Nikon). Construct area from n = 8–11 hydrogels per group from 2 HTM/GTM cell strains with three to four replicates per HTM/GTM cell strain was measured using FIJI software and normalized to 0 days followed by normalization to controls.

HTM Hydrogel Cell Proliferation Analysis

Cell proliferation was measured with the CellTiter 96® Aqueous Non-Radioactive Cell Proliferation Assay (Promega) following the manufacturer’s protocol. HTM hydrogels cultured in DMEM with 10% FBS and 1% PSG in presence of the different treatments for 5 days were incubated with the staining solution (38 µL MTS, 2 µL PMS solution, 200 µL DMEM) at 37°C for 1.5 h. Absorbance at 490 nm was recorded using a spectrophotometer plate reader (BioTEK, Winooski, VT, United States). Blank-subtracted absorbance values served as a direct measure of HTM cell proliferation from n = 6–8 hydrogels per group from 2 HTM/GTM cell strains with three to four replicates per HTM/GTM cell strain.
HTM Hydrogel Rheology Analysis

Fifty microliters of acellular hydrogel precursor solutions were pipetted into custom 8x1-mm PDMS molds. Similarly, 250 µL of HTM cell-laden hydrogel precursor solutions were pipetted into 16x1-mm PDMS molds. All samples were UV crosslinked and equilibrated as described above. Acellular hydrogels were measured at 0 days. HTM hydrogels, cultured in DMEM with 10% FBS and 1% PSG in presence of the different treatments, were measured at 5 days; samples were cut to size using an 8-mm diameter tissue punch. A Kinexus rheometer (Malvern Panalytical, Westborough, MA, United States) fitted with an 8-mm diameter parallel plate was used to measure hydrogel viscoelasticity. To ensure standard conditions across all experiments (n = 3 per group), the geometry was lowered into the hydrogels until a calibration normal force of 0.02 N was achieved. Subsequently, an oscillatory shear-strain sweep test (0.1–60%, 1.0 Hz, 25°C) was applied to determine storage modulus (G’), and loss modulus (G″) in the linear region. Elastic modulus was calculated with \( E = 2\times (1 + v)\times G' \), where a Poisson’s ratio \( v \) of 0.5 for the ECM hydrogels was assumed (Timothy and Lodge, 2020).

HTM Hydrogel Atomic Force Microscopy Analysis

Thirty microliters of HTM cell-laden hydrogel solution were pipetted onto a Sursasil (Fisher Scientific) coated 12-mm round glass coverslip followed by placing a regular 12-mm round glass coverslip onto the hydrogels. Constructs were crosslinked by exposure to UV light at 320–500 nm, 2.2 W/cm² for 5 s, and cultured in DMEM with 10% FBS and 1% PSG for 7 days. Samples were shipped overnight to UHCO where they were locally masked. Elastic modulus of HTM cell-laden hydrogels were obtained by atomic force microscopy (AFM) in fluid in contact mode as described previously (Raghunathan et al., 2018). Briefly, force vs. indentation curves were obtained on a Bruker BioScope Resolve (Bruker nanoSurfaces, Santa Barbara, CA) AFM employing a PNP-TR cantilever (NanoAndMore, Watsonville, CA) whose pyramidal tip was modified with a borosilicate bead (nominal diameter 5 µm), nominal spring constant of 0.32 N/m assuming sample Poisson’s ratio of 0.5. The diameter of each sphere attached to the cantilever was qualified prior to usage. Prior to experimental samples, cantilever was calibrated on a silicon wafer in fluid. Force-distance curves were randomly obtained from 7–10 locations with three force curves obtained per location per sample. Force curves were analyzed using the Hertz model for spherical indenters using a custom MATLAB code (Chang et al., 2014). Curve fits were performed for at least 1 µm of indentation depth for the samples.

Statistical Analysis

Individual sample sizes are specified in each figure caption. Comparisons between groups were assessed by unpaired t-test, one-way or two-way analysis of variance (ANOVA) with Tukey’s multiple comparisons post hoc tests, as appropriate. The significance level was set at \( p < 0.05 \) or lower. GraphPad Prism software v9.2 (GraphPad Software, La Jolla, CA, United States) was used for all analyses.

RESULTS

TGFβ2 and YAP/TAZ Activity is Upregulated in GTM Cells

It has been shown that levels of TGFβ2 are elevated in eyes of glaucomatous patients compared to age-matched normal eyes (Inatani et al., 2001; Picht et al., 2001; Ochiai and Ochiai, 2002; Agarwal et al., 2015). We confirmed that GTM cells isolated from donor eyes with POAG history secreted significantly more active TGFβ2 protein by ~2.44-fold (Figure 1A) compared to normal HTM cells, which was consistent with increased mRNA levels (Supplementary Figure S2A).

To investigate YAP and TAZ transcriptional activity under normal and glaucomatous conditions, the expression of YAP/TAZ and relevant select downstream targets at mRNA and protein levels were evaluated in normal HTM and GTM cells, which were plated with similar cell density. The functions of YAP/TAZ depend on their spatial localization within the cellular nucleus or cytoplasm. When localized to the nucleus, YAP/TAZ interact with TEAD transcription factors to drive the expression of certain proteins, such as transglutaminase-2 (TGM2) and connective tissue growth factor (CTGF), cysteine-rich angiogenic inducer 61 (CYR61) and ankyrin repeat domain 1 (ANKRD1) (Low et al., 2014). When localized in the cytoplasm, YAP can be phosphorylated at five serine/threonine residues, TAZ at four, by the large tumor suppressor (LATS) kinases 1 and 2. Of these sites, the most relevant residues that keep YAP and TAZ inhibited are S127 (S89 in TAZ) and S381 (S311 in TAZ) (Zhao et al., 2010). Phosphorylation of YAP/TAZ either primes to binding with 14-3-3σ leading to their cytoplasmic sequestration or ubiquitin-mediated protein degradation (Zhao et al., 2007; Lei et al., 2008; Dupont et al., 2011). We demonstrated that mRNA levels of YAP and TAZ were both significantly upregulated in GTM cells compared to normal HTM cells (Figure 1B). TGM2 and CTGF have been shown to play a role in HTM cell pathobiology in glaucoma (Tovar-Vidales et al., 2008; Junglas et al., 2012); here we showed that mRNA of TGM2, CTGF and ANKR1D1 were also significantly upregulated in GTM cells vs. normal HTM cells, whereas no significant difference was observed for CYR61 (Figure 1B, Supplementary Figures S2B,C). GTM cells showed significantly lower p-YAP (S127) and p-TCG (S89) vs. normal HTM cells, while total YAP and TAZ expression were equivalent to HTM cells; this resulted in significantly decreased p-YAP/YAP and p-TCG/TCG ratios (= less inactive YAP/TAZ in GTM cells). Consistent with the lower ratio of phosphorylated-to-total proteins, 14-3-3σ expression in GTM cells was significantly decreased compared to normal HTM cells (Figures 1C,D). Besides, GTM cells exhibited significantly increased YAP/TAZ nuclear-to-cytoplasmic (N/C) ratio and TGM2 expression compared to normal HTM cells (= more active YAP/TAZ in GTM cells) (Figures 1E–H, Supplementary Figure S3). Together, these data demonstrate that levels of active TGFβ2 as well as YAP and TAZ nuclear...
Localization and transcriptional activity are elevated in GTM cells isolated from patients with glaucoma compared to normal HTM cells.

ECM Stiffening Increases YAP/TAZ Activity in HTM Cells via Modulating Focal Adhesions and Cytoskeletal Rearrangement

Atomic force microscopy (AFM) analyses showed that the TM from POAG eyes is ~1.5-5-fold stiffer compared to that from healthy eyes (Wang et al., 2017a; Wang et al., 2017b; Vahabikashi et al., 2019). To that end, we recently showed that DEX treated HTM cell-laden hydrogels are ~2-fold stiffer compared to controls (Li et al., 2021a). Here, we demonstrated that TGFβ2-treated HTM cell encapsulated hydrogels were also ~2-fold stiffer compared to controls using AFM and rheology (Figure 2A, Supplementary Figure S4). To mimic the stiffness difference between glaucomatous and healthy HTM tissue, we utilized riboflavin (RF)-mediated secondary UV crosslinking of collagen fibrils, which stiffened the hydrogels by ~2-fold (Figure 2B).

Adherent cells are connected to the ECM through transmembrane receptor integrins and focal adhesion (FA) proteins such as vinculin (Mohammed et al., 2019). HTM cells on stiff glass exhibited strong vinculin FA staining (Supplementary Figure S5A), while cells on soft hydrogels showed qualitatively fewer vinculin puncta compared to cells on glass (Figure 2C). The stiffened ECM hydrogels significantly increased number and size of vinculin puncta in HTM cells compared to HTM cells on the soft hydrogel substrates (Figures 2C,D; Supplementary Figure S5B). Consistent with our observation on vinculin FA, HTM cells on stiff hydrogels exhibited ~1.26-fold larger nuclei compared to cells on the soft matrix, which was still less compared to the nuclei in HTM cells on conventional glass (Figures 2E,F). We also found that the stiffened ECM hydrogels significantly increased filamentous (F)-actin, phospho-myosin light chain (p-MLC) and α-smooth muscle actin (αSMA) levels in HTM cells compared to cells on the soft hydrogels, which was still less compared to the nuclei in HTM cells on conventional glass (Figures 2G–J, Supplementary Figures S5C,D). Importantly, we observed that nuclear localization of YAP/TAZ, TGM2 expression, and fibronectin (FN) remodeling in HTM cells were upregulated by the stiffened ECM hydrogels compared to cells on the soft hydrogels (Figure 2K–N; Supplementary Figures S5E–H).

Together, these results suggest that the stiffened ECM hydrogels induce a bigger nuclear size in HTM cells on stiff matrix compared to cells on soft matrix. The stiffened hydrogels induce FA and actomyosin cytoskeletal rearrangement,
correlating with elevated YAP/TAZ nuclear localization in HTM cells.

**TGFβ2 Regulates YAP/TAZ Activity via ERK and ROCK Signaling Pathways**

TGFβ2 has been shown to activate canonical Smad (Smad2/3) and diverse non-Smad signaling pathways including extracellular-signal-regulated kinase (ERK), c-Jun N-terminal kinases, P38 kinases and Rho-associated kinase (ROCK) in various cell types with context-dependent crosstalk (Zhang, 2009; Prendes et al., 2013; Montecchi-Palmer et al., 2017; Ma et al., 2020). In HTM cells, TGFβ2 was shown to upregulate ERK via activating RhoA/ROCK to induce a fibrotic-like, contractile cell phenotype using conventional stiff culture substrates (Pattabiraman and Rao, 2010). By contrast, we recently demonstrated that in the context of cell contractility, ROCK activity was negatively regulated by ERK with TGFβ2 induction in HTM cells cultured on soft tissue-like ECM hydrogels; this resulted in differentially regulated F-actin, αSMA, FN, and p-MLC (Li et al., 2021b). Here, we investigated whether YAP signaling in HTM cells in response to TGFβ2 induction requires ERK or ROCK. HTM or GTM cells were seeded on top of soft hydrogels treated with TGFβ2 ± U0126 (ERK inhibitor) or Y27632 (ROCK inhibitor). Consistent with our observation for cells seeded on glass (Figure 1), we found

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**FIGURE 2 | Stiffened ECM hydrogels elevate YAP/TAZ activity in HTM cells via modulating FA and cytoskeletal rearrangement.** (A) Elastic modulus of HTM cell-encapsulated hydrogels subjected to control and TGFβ2 (2.5 ng/ml) measured by AFM (n = 6 replicates/group). (B) Schematic of riboflavin (RF)-mediated double photo-crosslinking to stiffen ECM hydrogels, and elastic modulus of the hydrogels (n = 3 replicates/group). (C) Representative fluorescence micrographs and Fiji mask images of vinculin in HTM cells on soft and stiff hydrogels (dashed box shows region of interest in higher magnification image; vinculin = grey). Scale bar, 20 µm (left) and 5 µm (right). (D) Analysis of number of vinculin puncta (n = 20 images from 2 HTM cell strains with three replicates per cell strain). (E,F) Representative fluorescence micrographs and size analysis of HTM cell nuclei on soft and stiff hydrogels (n = 30 images from 2 HTM cell strains with 3–6 replicates per cell strain; more than 100 nuclei were analyzed per cell strain; dotted line shows mean nuclear area of HTM cells on glass for reference). (G,J) Representative fluorescence micrographs of F-actin and p-MLC in HTM cells on soft and stiff hydrogels (dashed box shows region of interest in higher magnification image. F-actin = green; p-MLC = red). Scale bar, 20 µm. (H,J) Analysis of F-actin and p-MLC intensity (n = 20 images per group from 2 HTM cell strains with three replicates per HTM cell strain). (K,M) Representative fluorescence micrographs of YAP/TAZ in HTM cells on soft and stiff hydrogels (YAP/TAZ = grey). Scale bar, 20 µm. (L,N) Analysis of YAP/TAZ nuclear/cytoplasmic ratio (n = 30 images from 2 HTM cell strains with 3–6 replicates per cell strain). Open and closed symbols, or symbols with different colors represent different cell strains. The lines and error bars indicate Mean ± SD; Significance was determined by unpaired t-test (**p < 0.01; ***p < 0.001; ****p < 0.0001).
significantly higher levels of nuclear YAP/TAZ and TGM2 expression in GTM cells compared to normal HTM cells on
the soft ECM hydrogels (Figure 3; Supplementary Figure S6). TGFβ2 increased nuclear YAP in both HTM and GTM cells, which were blocked by co-treatment of U0126 or Y27632. Importantly, we found that YAP N/C ratio in GTM cells with ERK or ROCK inhibition were at similar levels as HTM controls (Figures 3A–D).

YAP and TAZ are generally thought to function similarly in response to mechanical and biochemical signals (Totaro et al., 2018). However, it has been shown that YAP and TAZ are distinct effectors of TGFβ1-induced myofibroblast transformation (Muppala et al., 2019). In this study, we observed that TGFβ2 significantly induced nuclear TAZ localization in HTM cells, and co-treatment of U0126 or Y27632 with TGFβ2 rescued TAZ nuclear localization to control levels (Figures 3E,F). Similarly, TGFβ2 increased TAZ N/C ratio in GTM cells, while ERK or ROCK inhibition decreased TGFβ2-induced nuclear TAZ to HTM control levels (Figures 3G,H). Consistent with YAP/TAZ nuclear localization, TGM2 expression was upregulated by TGFβ2, which was significantly decreased by co-treatment of U0126 or Y27632 (Supplementary Figure S6).

These data show that TGFβ2 increases nuclear YAP/TAZ and TGM2 expression in both HTM and GTM cells, which was attenuated by either U0126 or Y27632 co-treatment. Collectively, this suggests that external biophysical (stiffened ECM) and biochemical signals (increased TGFβ2) drive altered YAP and TAZ mechanotransduction.

**FIGURE 3** Effects of TGFβ2 in absence or presence of ERK or ROCK inhibition on nuclear YAP/TAZ localization in HTM and GTM cells. (A,C,E,G) Representative fluorescence micrographs of YAP or TAZ in HTM or GTM cells on soft hydrogels subjected to control, TGFβ2 (2.5 ng/ml), TGFβ2 + U0126 (10 µM), TGFβ2 + Y27632 (10 µM) at 3 days (YAP = grey). Scale bar, 20 µm. (B,D,F,H) Analysis of YAP or TAZ nuclear/cytoplasmic ratio (n = 20 images from 2 HTM cell strains with three replicates per cell strain; n = 10 images from one GTM cell strain with three replicates). Symbols with different colors represent different cell strains; dotted line shows HTM cell control mean value for reference. The lines and error bars indicate Mean ± SD. Significance was determined by one-way ANOVA using multiple comparisons tests (shared significance indicator letters represent non-significant difference (p > 0.05), distinct letters represent significant difference (p < 0.05)).
in HTM cells that contributes to glaucomatous cellular dysfunction.

**F-Actin Polymerization Modulates YAP/TAZ Activity**

It has been shown that latrunculin B (Lat B), a compound that inhibits polymerization of the actin cytoskeleton, increases outflow facility and decreases IOP in human and non-human primate eyes (Peterson et al., 1999; Tian et al., 2000; Ethier et al., 2006). Likewise, Lat B has been reported to depolymerize HTM cell actin and decrease cellular stiffness via cytoskeletal relaxation (McKee et al., 2011). To investigate the effects of F-actin cytoskeletal disruption on YAP/TAZ activity, HTM cells were seeded on soft hydrogels, and treated with TGFβ2 for 3 days, followed by Lat B treatment for 30 min (Figure 4A). Consistent with our previous study (Li et al., 2021b), TGFβ2 significantly increased F-actin fibers, and co-treatment of Lat B with TGFβ2 potently depolymerized F-actin and decreased F-actin fiber formation (Figures 4B,C) without negatively influencing cell viability (all three groups exhibited ~8 cells/0.05 mm²). We observed significantly more vinculin puncta and increased puncta size induced by TGFβ2, which was abolished by Lat B treatment (Figures 4D,E; Supplementary Figure S7). Similar to before (Figure 3), TGFβ2 significantly increased YAP/TAZ nuclear localization and TGM2 expression, which were restored to control levels by 30 min of Lat B co-treatment (Figures 4F–K). Immunoblot analyses corroborated the immunostaining results and showed that co-treatment of TGFβ2 and Lat B increased the ratio of p-YAP to YAP and p-TAZ to TAZ (Figures 4L–M), representing overall decreased nuclear YAP and TAZ consistent with previous studies (Piccolo et al., 2014; Das et al., 2016; Panciera et al., 2017). Interestingly, according to the immunoblot result, Lat B may regulate YAP
activity by increasing p-YAP while maintaining similar levels of total YAP, whereas Lat B may decrease total levels of TAZ (Figures 4L,M).

In sum, these findings demonstrate that F-actin depolymerization decreases TGFβ2-induced vinculin FA, nuclear YAP/TAZ and their downstream target TGM2.

**YAP/TAZ Regulate FA Formation, ECM Remodeling and Cell Contractile Properties**

Our results so far suggest that POAG-related stimuli (i.e., TGFβ2 and stiffened ECM) increase YAP/TAZ activity in HTM cells. To further investigate the roles of YAP/TAZ in HTM cell function, we seeded HTM cells on stiff hydrogels to increase the baseline levels of YAP/TAZ in controls, and depleted YAP and TAZ using combined siRNA knockdown (Figure 5A). Treatment reduced mRNA expression levels to 34.52 and 35.25% of siRNA controls, respectively (Supplementary Figure S8A). On a protein level, we observed 64.80% YAP knockdown with combined siYAP/TAZ treatment or 49.10% YAP knockdown with single siYAP treatment. TAZ protein levels were reduced by 89.90% with siYAP/TAZ and by 80.90% with siTAZ treatments, respectively (Supplementary Figures S8B–D). Curiously, we noted a drop in TAZ protein levels in siYAP-treated HTM cells, whereas no such effects were noted in YAP protein expression with siTAZ treatment, requiring further investigation.

**FIGURE 5 | YAP/TAZ are regulators of FA formation, ECM remodeling and cell contractile properties in HTM cells.** (A) Schematic showing time course of YAP/TAZ depletion using siRNA experiments with HTM cells. (B,D,F,H,J and L) Representative fluorescence micrographs of TGM2, FN, aSMA, F-actin, p-MLC and vinculin in HTM cells on stiff hydrogels subjected to siControl or siYAP/TAZ (dashed box shows region of interest in higher magnification image; TGM2/F-actin = green; FN/p-MLC/aSMA = red; vinculin = grey; DAPI = blue). Scale bar, 20 μm in B,F,H,J; 20 and 5 μm in L; 100 μm in D. (C,E,G,I,K,M) Analysis of TGM2, FN, aSMA, F-actin, p-MLC and number of vinculin puncta (n = 20 images from 2 HTM cell strains with three replicates per cell strain). (N) Representative fluorescence micrographs and size analysis of HTM cell nuclei on stiff hydrogels subjected to siControl or siYAP/TAZ (n = 40 images from 2 HTM cell strains with six replicates per cell strain; more than 100 nuclei were analyzed per cell strain). (O) mRNA fold-change of FN, TGM2, CTGF, aSMA in HTM cells on stiff hydrogels subjected to siControl or siYAP/TAZ by qRT-PCR. The mRNA levels were normalized to the levels of GAPDH mRNA 48 h post transfection (n = 6 replicates from 2 HTM cell strain). Open and closed symbols indicate Mean ± SD; Significance was determined by unpaired t-test (C,E,G,I,K,M,N) and two-way ANOVA (O) using multiple comparisons tests (**p < 0.01; ***p < 0.0001).
investigation. Immunostaining showed that YAP/TAZ nuclear localization in HTM cells transfected with siYAP/TAZ were significantly decreased vs. controls despite culture in a stiffened ECM environment (Supplementary Figures S8E–H). Importantantly, we observed that YAP/TAZ depletion significantly reduced expression of TGM2, FN and CTGF, which may decrease HTM ECM stiffness (Figures 5B–E, O).

F-actin filaments, αSMA and p-MLC are all involved in cell contractility regulation, and we have previously demonstrated that their expression was upregulated in HTM cells under simulated glaucomatous conditions (Li et al., 2021a; Li et al., 2021b). Here, YAP/TAZ depletion significantly decreased F-actin filaments and expression of αSMA and p-MLC, suggestive of reduced HTM cell contractility (Figures 5F–I, O). We also found that the expression of canonical Hippo pathway kinases LATS1, LATS2 and 14-3-3σ was not affected by siYAP/TAZ knockdown (Supplementary Figure S8I). Tension generated within the actomyosin cytoskeleton is transmitted across FA to induce integrin-mediated remodeling of the ECM (Jansen et al., 2017). As such, we observed that YAP/TAZ depletion significantly decreased the number and size of vinculin puncta and nuclear size (Figures 5L–N; Supplementary Figure S8J).

Collectively, these data demonstrate that YAP/TAZ depletion using siRNA leads to impaired YAP/TAZ signaling, FA formation, cytoskeletal/nuclear and ECM remodeling, and cell contractile properties.

**YAP/TAZ-TEAD Interaction is Required for YAP/TAZ Downstream Effects**

Canonically, YAP/TAZ bind to TEAD family members to induce the transcription of YAP/TAZ target genes (Low et al., 2014). To test the effects of YAP/TAZ-TEAD interaction on HTM cell behavior under simulated glaucomatous conditions, we tested the effects of verteporfin (VP), a selective inhibitor of the YAP/TAZ-TEAD transcriptional complex (Liu-Chittenden et al., 2012). HTM cells were seeded on top of or encapsulated in soft hydrogels, and treated with TGFβ2 ± VP. Interestingly, we observed that HTM cells inside hydrogels showed qualitatively decreased YAP nuclear-to-cytoplasmic ratio compared to cells atop of the hydrogels in absence of any treatments (Figures 6A–D). In both cell culture environments, TGFβ2 increased nuclear YAP/TAZ localization, and treating HTM cells with VP significantly decreased TGFβ2-induced nuclear YAP/TAZ as well as downstream TGM2 compared to controls, consistent with the effects of siRNA-mediated YAP/TAZ depletion (Figures 6A–D; Supplementary Figures S9A–D). We observed that the inhibition of YAP/TAZ-TEAD interaction not only decreased nuclear YAP/TAZ, but also reduced cytoplasmic and total protein

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**FIGURE 6** | Inhibition of YAP/TAZ-TEAD interaction with VP decreases nuclear YAP and F-actin levels. (A) Representative fluorescence micrographs of YAP in HTM cells cultured on top of soft hydrogels subjected to control, TGFβ2 (2.5 ng/ml), VP (0.5 µM), TGFβ2 + VP at 3 days (YAP = grey; DAPI = blue). Scale bar, 20 µm. (B) Analysis of YAP nuclear/cytoplasmic ratio (n = 20 images from 2 HTM cell strains with three replicates per cell strain). (C) Representative fluorescence micrographs of YAP in HTM cells encapsulated in soft hydrogels subjected to the different treatments at 3 days (YAP = grey; DAPI = blue). Arrows indicate higher YAP in nuclei, arrowheads indicate higher YAP in cytoplasm. Scale bar, 20 µm. (D) Analysis of YAP nuclear/cytoplasmic ratio (n = 6 images from one HTM cell strain with three replicates). (E) Representative fluorescence micrographs of F-actin in HTM cells on or inside of soft hydrogels subjected to the different treatments (F-actin = red). Scale bar, 50 µm. (F) Analysis of F-actin intensity in HTM cells on soft hydrogels (n = 20 images from 2 HTM cell strains with three replicates per cell strain). (G) Analysis of F-actin intensity in HTM cells inside of soft hydrogels (n = 6 images from one HTM cell strain with three replicates). Open and closed symbols represent different cell strains. The lines and error bars indicate Mean ± SD; Significance was determined by one-way ANOVA using multiple comparisons tests [shared significance indicator letters represent non-significant difference (p > 0.05), distinct letters represent significant difference (p < 0.05)].
levels ([Supplementary Figure S9E](#)). VP treatment significantly decreased TGFβ2-induced F-actin filaments, expression of αSMA and p-MLC, and FN deposition ([Figures 6E–G; Supplementary Figures S9F–K](#)).

Collectively, these results show that pharmacological inhibition of YAP/TAZ-TEAD interaction leads to impaired ECM remodeling and cell contractile properties, which is independent of cell culture environments (atop or encapsulated in ECM hydrogels).

**YAP/TAZ Mediate HTM Cell Contractility and HTM Hydrogel Stiffness**

Lastly, to investigate the effects of YAP/TAZ on HTM cell contractility and ECM stiffening, we encapsulated HTM cells in ECM hydrogels and treated with TGFβ2, either alone or in combination with VP, and assessed hydrogel contractility and stiffness. TGFβ2-treated HTM hydrogels exhibited significantly greater contraction vs. controls by 5 days (74.30% of controls), consistent with our previous report ([Li et al., 2021a](#)). Co-treatment of TGFβ2 + VP potently decreased HTM hydrogel contraction (90.37% of controls) compared to TGFβ2-treated samples ([Figure 7A](#)). We observed that TGFβ2 significantly increased hydrogel stiffness (1.86-fold of controls), which was prevented by co-treatment with VP (1.16-fold of controls) ([Figure 7B](#)).

To assess whether YAP/TAZ inhibition had comparable effects on GTM cells, we evaluated GTM cell-laden hydrogel contraction in response to the same treatments. Consistent with our previous study ([Li et al., 2021a](#)), we demonstrated that GTM hydrogels in absence of additional TGFβ2 induction exhibited significantly greater contraction relative to normal HTM hydrogels (86.08% of HTM hydrogels controls); TGFβ2 further increased GTM hydrogel contraction, and VP partially blocked this during the short 5 days exposure ([Figure 7C](#)).

To determine if hydrogel contractility was influenced by the cell number, we assessed HTM/GTM cell proliferation in constructs subjected to the different treatments. We observed a fewer number of cells in the TGFβ2 + VP group compared to TGFβ2-treated samples (10.10% decreased) ([Supplementary Figure S10A](#)), while co-treatment of TGFβ2 + VP reduced hydrogel contraction and stiffening by 21.63 and 59.45% compared to the TGFβ2-treated group, respectively ([Figures 7A,B](#)); demonstrating that VP induced decreasing hydrogel contraction was not only caused by the smaller cell number, but also reduced cell contractility. No significant differences between the different groups were observed for GTM cell-laden hydrogels ([Supplementary Figure S10B](#)).
Together, these data demonstrate that TGFβ2 robustly induces HTM hydrogel contractility and stiffening in a soft ECM environment, which are potently reduced by YAP/TAZ inhibition. Likewise, inhibition of YAP/TAZ had similar effects on GTM cells inside the 3D hydrogel network.

DISCUSSION

The mechanosensitive transcriptional coactivators YAP and TAZ play important roles in mechanotransduction, a process through which cells translate external biophysical cues into internal biochemical signals. YAP/TAZ modulate target gene expression profiles with broad functional consequences across many cell and tissue types (Dupont et al., 2011; Boopathy and Hong, 2019). Through this mechanism, YAP/TAZ signaling regulates critical cellular functions and normal tissue homeostasis; imbalance or failure of this process is at the core of various diseases (Panciera et al., 2017). Indeed, elevated YAP/TAZ transcriptional activity is associated with glaucomatous HTM cell dysfunction (Thomasy et al., 2013; Chen et al., 2015; Ho et al., 2018; Peng et al., 2018; Dhamodaran et al., 2020; Yemanyi et al., 2020; Yemanyi and Raghunathan, 2020). Importantly, a recent genome-wide meta-analysis identified YAP among 44 previously unknown POAG risk loci (Gharahkhani et al., 2021); this observation provides strong new evidence that YAP may play a prominent role in glaucoma pathogenesis. Here, we found that YAP/TAZ nuclear localization, the principal mechanism to regulate their function, is significantly higher in TM cells from patients with glaucoma compared to cells isolated from healthy tissue, exhibiting normal variability between cells from different donors (i.e., one GTM cell strain showed higher baseline YAP/TAZ nuclear localization and transcriptional activity compared to the other) (Figure 1; Supplementary Figures S2, S3). However, the detailed mechanisms for YAP/TAZ modulation in HTM cells under glaucomatous conditions (i.e., stiffened ECM and increased growth factors in AH) remain to be elucidated. To model this, we used biomimetic ECM hydrogels with tunable stiffness to study the roles of YAP and TAZ in HTM cells in response to stiffened matrix and TGFβ2. As summarized in Figure 8, our data support that YAP/TAZ are critical regulators in mediating HTM cellular responses to the stiffened ECM and elevated TGFβ2 in POAG involving both ERK and ROCK signaling with likely crosstalk; we propose that increased YAP/TAZ nuclear retention may drive further HTM tissue stiffening to exacerbate disease pathology conditions. This conclusion is supported by the findings that (i) stiffened ECM elevates YAP/TAZ nuclear localization, potentially through regulating FA formation and actin cytoskeleton rearrangement; (ii) TGFβ2 induces nuclear YAP/TAZ localization and target gene activation through ERK and ROCK signaling pathways; (iii) depolymerization of F-actin decreases nuclear YAP/TAZ; (iv) YAP/TAZ depletion using siRNA or pharmacological inhibition of YAP/TAZ-TEAD
interaction demonstrated that TGFβ inhibitor VP decreases HTM/GTM cell-laden hydrogel contraction and HTM stiffening.

Most in vitro studies of HTM cell (patho-)physiology have relied on conventional cell monolayer cultures on plastic or glass of supraphysiologic stiffness. However, biophysical cues such as substrate composition and stiffness are known to be potent modulators of cell behaviors (Raghunathan et al., 2013). In our previous studies, we reported that HTM cells on glass exhibited bigger nuclei and different organization of F-actin fibers compared to HTM cells on ECM hydrogels (Li et al., 2021b). Here, we demonstrated that HTM cells on hydrogels showed a smaller number of vinculin FA vs. HTM cells on glass (Figures 2C,D; Supplementary Figure S5A), confirming that HTM cells display distinct physiological characteristics on soft tissue-mimetic biomaterials and traditional tissue culture plastic or glass substrates of supraphysiologic stiffness (Caliani and Burdick, 2016).

Increased tissue stiffness has been observed in multiple pathologies, including cancer, cardiovascular and fibrosis-related diseases (Lampi and Reinhart-King, 2018). ECM stiffening can precede disease development and consequently increased mechanical cues can drive their progression via altered mechanotransduction (Frantz et al., 2010; Kaess et al., 2012; Pickup et al., 2014). Therefore, therapeutically targeting ECM stiffening by disrupting the cellular response to the stiffened ECM environment, or in other words targeting mechanotransduction, is an emerging field with clear implications for glaucoma treatment. It is widely accepted that the ECM is stiffer in the glaucomatous HTM (Wang et al., 2017b). Previous studies have also shown that ECM deposited by HTM cells treated with DEX was ~2-4 fold stiffer relative to controls (Raghunathan et al., 2015; Raghunathan et al., 2018). Our recent data using DEX-induced HTM cell-encapsulated 3D ECM hydrogels were in good agreement with these observations (Li et al., 2021a). In this study, we demonstrated that TGFβ2 treatment resulted in a comparable ~2-fold increase in ECM stiffness surrounding the HTM cells as they reside embedded in our bioengineered hydrogels (Figure 2A). In both scenarios, the stiffness changes were cell-driven in response to biochemical cues implicated in glaucoma.

Corneal UV crosslinking with riboavin (RF) is a clinical treatment to stabilize the collagen-rich stroma in corneal ectasias (Zhang et al., 2011; Rahman et al., 2020), with promising potential for enhancing mechanical properties of collagen-based hydrogels (Ahearne and Coyle, 2016; Heo et al., 2016). Crosslinking occurs via covalent bond formation between amine acids of collagen fibrils induced by singlet O2 from UV-excited RF (Tirella et al., 2012). To simulate glaucomatous ECM stiffening for investigations of the cellular response, we used roboavin to double-crosslink collagen in the hydrogels, which increased their stiffness ~2-fold over baseline (Figure 2B). HTM cells on the stiffened hydrogels exhibited bigger nuclei, increased number of vinculin FA, cytoskeleton rearrangement, ECM deposition and YAP/TAZ nuclear localization compared to cells on soft hydrogels (Figures 2C–N; Supplementary Figure S3). Taken together, our findings indicate that YAP/TAZ subcellular localization in HTM cells is distinctive on different substrates, suggesting that hydrogels with tunable physiochemical properties may be more suitable to investigate subtleties in HTM cell behaviors that would otherwise go unnoticed when relying exclusively on traditional stiff 2D culture substrates.

TGFβ2 levels are elevated in the aqueous humor of glaucoma patients compared to age-matched normal eyes (Inatani et al., 2001; Picht et al., 2001; Ochiai and Ochiai, 2002; Agarwal et al., 2015). Here, we showed that GTM cells isolated from POAG donor eyes secreted significantly more active TGFβ2 compared to normal HTM cells (Figure 1A; Supplementary Figure S2). We observed that TGFβ2 upregulated YAP/TAZ nuclear localization and TGM2 expression, a downstream effector of active YAP/TAZ signaling, in both normal HTM and GTM cells, confirming that YAP/TAZ activity are upregulated under glaucomatous conditions. Consistent with these observations, our immunoblot data showed significantly decreased p-YAP/YAP and p-TAZ/TAZ ratios (Figures 1C,D), indicative of “less inactive YAP/TAZ” in GTM cells compared to normal HTM cells. Given that S127 in YAP (and S89 in TAZ) is among the most relevant residues that keeps YAP inhibited, it appears likely that LATS1/2 kinases play a central role in HTM/GTM cell modulation. However, it is conceivable that kinases other than LATS1/2 target S127, as it was originally identified as AKT target (Basu et al., 2003). S127 phosphorylation has been observed to occur in a LATS1/2-independent manner; in the context of cell contractility and ECM stiffening, YAP mechanotransduction was also shown to occur independently of LATS-mediated phosphorylation, rather involving ROCK and myosin activities (Piccolo et al., 2014; Sorrentino et al., 2014).

ERK or ROCK inhibition decreased nuclear YAP/TAZ and TGM2 in both HTM and GTM cells, and co-treatment of ERK or ROCK inhibitor with TGFβ2 restored YAP/TAZ cellular localization and TGM2 to HTM control levels (Figure 3; Supplementary Figure S6). TGFβ2 is known to signal via both canonical Smad and non-canonical signaling pathways, such as ERK, JNK, P38 kinases and ROCK in various cell types (Zhang, 2009). In HTM cells, TGFβ2 induces crosslinked actin network formation and this process is similarly blocked by inhibition of Smad or non-Smad signaling pathways (Montecchi-Palmer et al., 2017). Among the non-canonical TGFβ2 signaling pathways, ERK and ROCK have emerged as potent regulators of F-actin, αSMA and FN expression - all of which are implicated in glaucomatous cell pathobiology (Pattabiraman and Rao, 2010; Montecchi-Palmer et al., 2017; Faralli et al., 2019). Other reports showed that ROCK promotes smooth muscle cell migration and serotonin-mediated cell proliferation via increasing ERK activity (Liu et al., 2004; Zhao et al., 2012). In neurons and microglia, ERK activity has been found to be negatively regulated by ROCK (Hensel et al., 2015). Together, this suggests the presence of complex ERK and ROCK signaling crosstalk to regulate cellular behaviors in a context-dependent manner; yet, the molecular mechanisms of their crosstalk in HTM cells is far from clear. TGFβ2 was shown to upregulate ERK via activating RhoA/ROCK in HTM cells cultured on conventional stiff substrates, such as glass or plastic, to induce a fibrotic-like, contractile cell phenotype (Pattabiraman et al., 2016).
and Rao, 2010). Consistent with other cell systems, findings from this study suggested that the two non-canonical signaling pathways act in sequence. In our recent study, however, we demonstrated that ERK may inhibit ROCK activity and p-MLC to increase contractility of HTM cells cultured in a soft tissue-like ECM environment; no such effect was observed in cells on stiff glass (Li et al., 2021b). Further research will be necessary to investigate in greater detail whether/how canonical Smad signaling cross talks with non-canonical ERK and ROCK signaling pathways to regulate YAP and TAZ activity in HTM cells.

Actomyosin cell contractility forces are increased in response to elevated ECM stiffness and TGFβ2 induction (Lampi and Reinhart-King, 2018; Li et al., 2021b). Additionally, we have demonstrated that the stiffened ECM induces YAP/TAZ nuclear localization, and increases F-actin filaments, p-MLC and αSMA - all involved in actomyosin cell contractility force generation. We hypothesized that increased actomyosin cell contractility may drive YAP/TAZ nuclear localization in HTM cells. Here, we used Lat B to depolymerize F-actin stress fibers and increase cytoskeletal relaxation. We found that a short exposure time to Lat B eliminated vinculin FA formation and YAP/TAZ nuclear localization, and decreased TGM2 expression. ROCK inhibition has also been shown to decrease HTM cell F-actin fibers and contractility (Li et al., 2021a). These observations on effects of Lat B were consistent with our findings that ROCK inhibitor reduced YAP/TAZ activity (Figures 3, 4F–M–M4F–M). It would be worthwhile to further investigate effects of other cell contractility related molecules, such as myosin light chain kinase, myosin II, cofillin and gelsolin, on YAP/TAZ activity in both HTM and GTM cells.

We have observed that simulated glaucomatous conditions (i.e., elevated TGFβ2 and stiffened ECM) upregulated nuclear YAP/TAZ, and treatments that can reduce IOP in experimental models (i.e., ROCK inhibitor and Lat B) downregulated YAP/TAZ nuclear localization, in agreement with previous reports (Piccolo et al., 2014; Das et al., 2016; Panciera et al., 2017). These observations led us to explore the role of YAP/TAZ on glaucoma pathology development. We found that YAP/TAZ depletion using siRNA consistently decreased FA formation (i.e., vinculin), and reduced expression of cell contractile proteins (i.e., F-actin, p-MLC and αSMA) and ECM proteins (i.e., FN) (Figure 5). Also, YAP/TAZ deactivation reduced expression of TGM2, a protein that promotes cell-matrix interactions and FN crosslinking to stiffen the ECM (Akimov et al., 2000), and CTGF, known to increase ECM production and cell contractility in HTM cells, and elevate IOP in mouse eyes (Junglas et al., 2012). Thus, YAP/TAZ inhibition may decrease subsequent ECM stiffness potentially through regulation of ECM, TGM2 and CTGF production. Furthermore, we demonstrated that inhibition of YAP/TAZ-TEAD interaction using VP significantly decreased TGFβ2-induced YAP/TAZ nuclear localization in HTM cells independent of their spatial arrangement atop or inside of the ECM hydrogels. Consistent with the effects of siRNA-mediated YAP/TAZ depletion, VP treatment rescued TGFβ2-induced YAP/TAZ activity, cell contractile properties and ECM remodeling (Figure 6; Supplementary Figure S9).

Notably, YAP/TAZ inhibition using VP blunted HTM/GTM cell-laden hydrogel contraction and stiffening (Figure 7). Thus, we conclude that YAP/TAZ act as central players in regulating HTM cell mechanical homeostasis in response to changes of the surrounding microenvironment (e.g., levels of growth factors, ECM stiffness) to maintain tissue-level structural integrity and functionality. It has been demonstrated that the relative roles of YAP/TAZ are cell type- and context-dependent; they can cause homeostatic regulation of tissue properties (negative feedback loop) or promote fibrotic conditions (positive feedback loop). Some reports implicated YAP/TAZ in a feed-forward promotion of cytoskeletal tension and ECM protein deposition (Lin et al., 2017; Nardone et al., 2017; Yemanyi and Raghunathan, 2020), while some research showed YAP/TAZ had a negative feedback regulation that acted to suppress actin polymerization and cytoskeletal tension (Qiao et al., 2017; Mason et al., 2019). Our data were consistent with the former; i.e., YAP/TAZ drives HTM cell contraction and ECM stiffening.

In conclusion, using our bioengineered tissue-mimetic ECM hydrogel system, we demonstrated that nuclear YAP/TAZ is upregulated in response to simulated glaucomatous conditions (i.e., TGFβ2 induction and stiffened ECM), and that YAP/TAZ activation induces HTM cell contractility and ECM remodeling, which together may increase HTM stiffness in POAG. Our findings provide strong evidence for a pathologic role of aberrant YAP/TAZ signaling in glaucomatous HTM cell dysfunction, and may help inform strategies for the development of novel multifactorial approaches to prevent progressive ocular hypertension in glaucoma.

DATA AVAILABILITY STATEMENT

All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

AUTHOR CONTRIBUTIONS

HL, VR, WDS, PSG, and SH designed all experiments, collected, analyzed, and interpreted the data. WDS provided the GTM cells. VR performed the AFM experiments. HL and SH wrote the manuscript. All authors commented on and approved the final manuscript. PSG and SH conceived and supervised the research.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcell.2022.844342/full#supplementary-material

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