The lens is the pivotal tissue in accommodation—the process by which the eye alters its focal distance from far to near. The lens continues to grow in size throughout a person’s lifetime while the size of the globe of the eye stays constant through adulthood. This growth is a result of lens epithelial cell (LEC) proliferation, which ultimately leads to an increase in the number of fiber cells. Since a lens fiber cell has a much larger volume than an LEC, the lens progressively becomes larger as a result of LEC proliferation. The age-related changes in lens size and shape contribute to presbyopia and cataracts.

The driving force(s) for this continuous growth remain unknown, at least in part due to technical challenges with reproducing the lens’ complex in vivo environment which includes biochemical and biomechanical influences. Flat-mount lens explants are frequently used to probe specific aspects of LEC behavior, but may not be appropriate for examining mechanobiological behavior due to altered cell morphology and mechanical environment. Similarly, species which accommodate using a human-like mechanism are few and prohibitive expensive for many basic scientific studies.

In disaccommodation, tension is applied to the lens capsule via the zonules due to relaxation of the ciliary muscle. During accommodation, this tension is released via ciliary muscle contraction, allowing the lens to elastically recoil to a rounder shape. The ciliary muscle remains active into old age and the mechanical properties of zonules are independent of age, implying that the human lens capsule may experience cyclic tension even after the onset of presbyopia. In the context of presbyopia, younger lenses exhibit a large magnitude in focal length changes during stretching, while lenses above the age of 60 showed no changes in focal length with stretching, such that the eye is only able to focus on distant objects. Axial strains acting upon the lens, similar to those experienced during disaccommodation, have been observed to lead to a reversible increase in LEC area indicating that mechanical loading of the lens is transduced onto its cells. This change in LEC area provides strong evidence that a lifetime of accommodation and disaccommodation and the resulting changes in LEC area would play a part in driving strain-responsive behaviors in LECs.

Recent work has demonstrated that the lens epithelium is sensitive to changes in its mechanical environment. LEC proliferation is altered during cataract surgery which disrupts both the biochemical and biomechanical homeostasis of the lens. Departure from equibiaxial stresses in the capsule may
drive cell migration and morphological changes leading to posterior capsular opacification (PCO). Such anisotropic strains have been found to exist in the intact human lens capsule near the equator, coinciding with the region in which proliferation is known to occur in the mouse lens. PO can be inhibited in vitro by using pharmaceutical agents targeting the cytoskeleton; the cytoskeleton is known to convey information about mechanical stresses on a cell to the nucleus resulting in changes in protein expression. Together, these results suggest that LECs are strain-responsive cells (i.e., they can alter their behavior in response to mechanical cues from their environment). However, these studies have primarily focused on pathologic LEC differentiation leading to PCO and therefore did not examine whether proliferation rates or biomarker expression levels were directly influenced by mechanical stretching.

Activation of YAP/Taz signaling is a potential mechanism by which the axial loading of the lens by disaccommodation is transduced onto the LECs and upregulates proliferative activity. Earlier studies have observed the expression of YAP in rodent LEC explants and the role it plays in FGF-induced LEC proliferation. The role of YAP in the mechanoregulation of LEC stretch-induced proliferation is not well established; however, stretch-induced YAP activation and a resulting upregulation of cell proliferation has been observed in other cell types, including epithelial cells from other tissue types (i.e., mammary, lung, and skin epithelium). It is well established that YAP has a pivotal role in transducing mechanical cues from stretching into increased cell proliferation. This study seeks to provide evidence that this same behavior is exhibited by LECs, thus identifying a contributing factor to the continuous growth of the lens throughout life.

In order to study the effects of lens stretching on the proliferation of LECs without disturbing the complex microenvironment present within the lens, it was necessary to culture whole lens tissues under different loading conditions for an extended period of time ex vivo. For this purpose, a lens stretching bioreactor was developed. Lens stretchers have been employed in previous studies; however, they were primarily used to provide insight on the biomechanical and optomechanical properties of the lens and were unsuitable for use in the sterile environment of an incubator. The lens stretcher developed for this study was capable of applying both static and cyclic loading conditions on the lens ex vivo, while remaining autoclavable to ensure sterility.

This study was designed to determine whether the mechanical forces experienced by the lens during the process of accommodation contribute to lens growth by increasing the proliferation rate of LECs as well as to identify what role YAP signaling plays in the stretch-induced proliferation of LECs. By developing a lens organ culture bioreactor capable of reproducing the biochemical milieu and cyclic tension exerted on the human lens throughout life, we have overcome the technical challenges associated with studying the intact lens epithelium ex vivo. This technique allows detailed study of the mechanobiologic response of the lens epithelium to stretching ex vivo while closely mimicking that experienced by the human lens in vivo. Furthermore, agonists or inhibitors of specific molecular pathways may be used to elucidate the underlying mechanobiologic mechanisms involved.

**METHODS**

All animal tissues were used in accordance with institutionally approved protocols.

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**Lens Stretching Organ Culture**

Freshly enucleated porcine eyes were obtained from a local abattoir (Delaware Meats, Delaware, OH, USA). Extraocular tissue was removed and the whole globe was disinfected by submergence in 0.5% povidone-iodine (Sigma-Aldrich Corp., St. Louis, MO, USA) in PBS for 5 minutes, then transferred into PBS. The globe was removed from the PBS and partial-thickness incisions were made with a scalpel along the limbus and the equator. The cornea and the iris were carefully removed. The globe was bisected along the equatorial incision and the posterior half was discarded. The posterior portion of the vitreous was removed and the anterior portion of the vitreous was left attached to the anterior section of the eye. Eight radial cuts were made through the sclera spaced 45° apart to create eight flaps surrounding the lens. To ensure uniform stretching about the lens’ circumference a stapler was used to attach each of the scleral flaps to a 40-mm diameter silicone disk (McMaster-Carr, Elmhurst, IL, USA) with a 10-mm diameter hole in the center with the lens positioned in the central hole (Fig. 1).

The silicone disks were mounted onto a bespoke stretching ring device which attached to eight equally-spaced peripheral holes in the silicone disk. Lenses mounted in this way were submerged in prewarmed, serum-free medium 199 (M199) with Earle’s salts, L-glutamine, and sodium bicarbonate, supplemented with 0.1% bovine serum albumin, 100 IU/mL penicillin, 100 mg/mL streptomycin and 2.5 mg/mL Amphotericin B (Sigma-Aldrich Corp.). Some lenses were exposed to the Yes-associated protein (YAP) function inhibitor verteporfin (5 mM verteporfin, Sigma-Aldrich Corp.). Lenses were incubated at 37°C, 5% CO₂ for 24 hours. After 23 hours in culture, 0.1% of 10 mM EdU (5-ethyl-2-deoxyuridine; Thermo-Fisher, San Jose, CA, USA) in DMSO was added to the cell culture media for the remaining 1 hour.

Lenses were subjected to static stretching at 0% (control), 6%, and 12% strain (defined as percent change in the equatorial diameter of the lens). These strain amplitudes were chosen so as to remain close to the in vivo physiological range. For lenses undergoing cyclic stretching, the stretching ring was mounted onto a motorized rig and stretched at 6% strain amplitude with a frequency of 0 (control), 0.05, 0.1, and 0.2 Hz. These frequencies were chosen with 0.2 Hz as the highest because previous studies have observed proliferation in other cell types is inhibited at higher frequencies. The strain amplitude was validated by comparing images of the lenses in the stretched and unstretched configuration in ImageJ (NIH, Bethesda, MD, USA).

**Flow Cytometry**

Lenses were removed from culture, immediately isolated from the surrounding tissue, and rinsed in PBS. The lens capsule was isolated by peeling the capsule open from the posterior pole using jeweler’s forceps then the fiber cell bundle removed. The lens capsules were rinsed twice with PBS and submerged in 0.25% trypsin with 0.04% ethylenediaminetetraacetic acid (EDTA) (VWR, Radnor, PA, USA) at 4°C for 18 hours. Excess trypsin was removed and the capsules were placed in a 57°C water bath for 30 minutes. We added 10 mL of M199 supplemented with 0.1% BSA to quench the trypsin activity. Capsule fragments were filtered using a 70-μm cell strainer and the solution was centrifuged for 10 minutes at 180 rcf to collect the LECs.

The LECs were immediately fixed in 10% neutral buffered formalin (Sigma-Aldrich Corp.) and stained for EdU detection with AlexaFluor 488-azide using a commercial kit for 30
minutes according to manufacturer instructions (Click-iT; Invitrogen, Carlsbad, CA, USA). LECs were also stained for nuclear detection using a commercial reagent for 30 minutes (NucRed Live 647 ReadyProbes; Invitrogen). LECs were analyzed using a BD LSR II flow cytometer using the 488- and 640-nm lasers (BD Biosciences, Franklin Lakes, NJ, USA).

The labeling index was calculated from the percentage of cells that had a positive signal for the AlexaFluor 488 stain using commercial software (Flowjo Turku, Finland).

The resulting raw data were analyzed by collecting measurements from unstained cells to establish autofluorescence thresholds and cells from the same lens stained with both dyes. The dataset from each stained sample was gated such that the threshold would exclude 95% of the signal from the unstained control.

**Fluorescent Microscopy**

In order to image the intact LEC monolayer in situ, a flat-mounting technique was used rather than imaging an intact lens. This avoids the potential complications arising from imaging a curved surface. Lenses were removed from culture, and immediate isolated from surrounding tissue. The intact lens was fixed in 10% neutral buffered formalin for 10 minutes in order prevent sloughing off of the LEC monolayer during flat-mounting. Lenses were then dissected and flat-mounts of the lens capsule were prepared. Lens flat-mounts were then fixed in 10% neutral buffered formalin for a further 10 minutes. The LECs were permeabilized in 0.05% (vol/vol) Triton X-100 and blocked in 1% (wt/vol) bovine serum albumin for 30 minutes. Flat-mounts were rinsed in PBS and then incubated with fluorescent stains. AlexaFluor 488-azide with a commer-
cial kit (Invitrogen) were used to visualize proliferative activity in the lens. To visualize YAP localization, flat mounts were incubated with primary and secondary antibodies for 60 minutes each. Primary antibodies included YAP1 Rabbit Polyclonal Antibody (Thermo Fisher Scientific). The secondary antibody used was Goat anti-Rabbit, AlexaFluor 488 (Thermo Fisher Scientific). For both tests, following incubation with the antibodies and commercially available kits, samples were then counterstained with Hoechst (Thermo Fisher Scientific). Lenses stained for proliferation were imaged using a confocal microscope (Nikon Eclipse Ti2-E; Nikon Instruments Inc., Melville, NY, USA) and those stained for YAP localization were imaged using a fluorescent microscope (Nikon Eclipse Ts2; Nikon Instruments Inc.).

Confocal images were processed to objectively remove background signal, assumed to arise from either autofluorescence or residual dye, as follows. Z-stacks were flattened by keeping the maximum channel intensity value for each pixel location in the horizontal plane. Pixels corresponding to Hoechst-positive nuclei were determined using Otsu’s threshold method to produce a binary mask. Morphologic closing and opening operations were applied to avoid the loss of real connections or development of spurious connections between pixels. Areas with an area >50 pixels were excluded. EdU-positive nuclei were detected as follows. Autofluorescence or residual dye intensity in the green channel was estimated on the basis of the green channel intensity for all pixels not corresponding to a nucleus (as defined above). An empirical (Kaplan-Meier) cumulative distribution function was determined by including all such pixels. A probability that the mean pixel intensity within a given nucleus was not a result of autofluorescence or residual dyes was then calculated. If this probability exceeded 95%, the corresponding nucleus was considered to be EdU-positive.

Statistical Analysis

A paired t test was conducted on paired lenses cultured for either 1 or 24 hours to determine whether the labeling indices of LECs, defined as the number of EdU-labeled LECs to the total number of LECs, varied with respect to time spent in culture. Simple linear regression analysis was used to determine the effects of stretch amplitude and frequency on LEC proliferation. Analysis of covariance (ANCOVA) was performed to investigate the effects of verteporfin on LEC proliferation across different stretching conditions. A post-hoc Tukey’s honestly significant difference (HSD) test was used to compare the verteporfin group with the control. Statistical analysis was performed using JMP Pro 13 (SAS Institute, Cary, SC, USA).

RESULTS

Labeling Index of LECs Varies with Culture Time

The microenvironment LECs experience within the eye in vivo differs from that experienced in vitro. In order to investigate whether the stresses caused by a change in environment affect the labeling index of LECs, paired whole lenses mounted on silicon rings and were cultured for either 1 hour or 24 hours, under null stretch conditions with each lens exposed to EdU for 1 hour. LECs were then analyzed using flow cytometry to determine labeling index.

A paired t test was used to determine if the labeling index of LECs cultured for 1 hour immediately following dissection was significantly different than those cultured for 24 hours (Fig. 2). A significant difference was found \( (P = 0.0055) \). The data show that the labeling index of LECs in whole lens cultures decreased with time in culture. The higher initial labeling index followed by a decrease over time may be due to a stress response induced by changing the LEC microenvironment and the subsequent acclimation of the LECs to the new environment. Longer culture times could therefore be preferable for later studies in order to avoid any initial confounding cellular response to a new microenvironment.

Increasing Stretch Amplitude Increases LEC Proliferation in Whole Lens Cultures

To determine if the amplitude of static stretching affected LEC proliferation, a total of eight pairs of whole lenses were cultured for a period of 24 hours under various static stretch conditions. For each pair, one was subjected to 6% (4 pairs) or 12% (4 pairs) strain while the other (control) was held at 0%. During the final hour of the culture period the lenses were exposed to EdU, which would be incorporated into any newly synthesized DNA. The LECs were analyzed using flow cytometry and the labeling index was calculated from the percentage of the total population of cells that had synthesized new DNA during the hour-long EdU pulse. The labeling index values for the different stretch conditions are presented in Table 1.

Simple linear regression analysis was used to determine if stretch amplitude was a significant predictor of LEC proliferation (Fig. 3). A significant regression equation was found \( \text{Labeling Index} = 1.03\% + 1.14\% \times \text{Percent Stretch} \), \( R^2 = 0.863, P < 0.0001 \). The data show that the proliferation of LECs increased proportionally to stretch amplitude. These findings indicate LEC proliferation is driven, at least in part, by mechanotransduction. Therefore, the accommodative process may contribute to the growth of the lens.

| Table 1. Labeling Indices of Lenses Cultured Under Varying Static Strain Amplitudes |
|---------------------------------|---------------------------------|-----|
| Strain Amplitude, %         | Labeling Index, %               | \( n \) |
| 0                             | 1.14 ± 0.37                     | 8   |
| 6                             | 3.25 ± 0.27                     | 4   |
| 12                            | 6.57 ± 0.46                     | 4   |
LEC Proliferation Increases with Stretching Frequency in Whole Lens Cultures

Once a link between LEC proliferation and stretch amplitude was established it was necessary to determine if a change in labeling index occurred in response to changes in stretching frequency as well as stretch amplitude. Whole lens tissues were cultured and analyzed as described above. Stretch amplitude was held at 6% and the triangular stretch waveform was applied cyclically, oscillating from 0% to 6%, at frequencies of 0, 0.05, 0.1, and 0.2 Hz. A total of 16 unpaired lenses were used. The labeling index values are described in Table 2.

Linear regression analysis was used to determine if LEC labeling index is predicted by changes in stretching frequency. Linear regression analysis predicted the relationship to follow: Labeling Index \(= 1.05 + 0.44 \times \text{Percent Strain Amplitude} \) \((R^2 = 0.965, P < 0.0001)\). Lenses were cultured for 24 hours under varying static strain amplitudes and exposed to a one-hour EdU pulse before LECs were isolated and analyzed using flow cytometry. The labeling index increased proportionally with strain amplitude, suggesting a strong relationship between lens stretching and LEC proliferation.

Stretching Alters the Localization of LEC Proliferation Across the Lens Capsule

Qualitative analysis of the effects of different stretching regimes on the localization of LEC proliferative activity was performed by staining flat-mounted lens capsules for the thymine analog, EdU, and counterstaining LEC with the nuclear stain Hoechst. Representative images of lenses cultured under null strain (Fig. 5A), 12% static stretch (Fig. 5B), and cyclic stretch at 6% amplitude and 0.20 Hz (Fig. 5C) were used (Fig. 5). Low levels of EdU staining were observed in the null stretch lens (Fig. 5A), primarily near the equator. The static stretch lens appeared to have a higher labeling index (Fig. 5B), with the majority of EdU staining also occurring near the equator. In the cyclic lens, EdU labeling was observed near the anterior pole (Fig. 5C) and at different points along the equator (Fig. 5D–F).

Inhibition of YAP Function Blocks Mechanotransductive Effects of Stretching on LEC Proliferation

The effects of the YAP function inhibitor verteporfin on LEC proliferation was determined by exposing both paired eyes to identical static stretching conditions: 0%, 6%, or 12%. Both members of each pair were cultured in enhanced M199 and the treatment group was supplemented with verteporfin. A total of 10 pairs of eyes were used with 4 pairs of eyes cultured under null stretch, and three pairs each with 6% and 12% static strain. The labeling index values are described in Table 3.

Linear regression analysis was performed on both treatment groups. A significant correlation was found for the control group (Labeling Index \(= 1.28 + 0.39 \times \text{Stretch Amplitude} \) \((R^2 = 0.975, P < 0.0001)\), but no significant relationship was found for the group treated with verteporfin (Labeling Index \(= 0.99 + 0.02 \times \text{Stretch Amplitude} \) \((R^2 = 0.292, P = 0.107; \text{Fig. 6})\). To confirm the difference between the treatment groups, ANCOVA was used to determine whether there was a significant difference between groups; a significant difference was observed \((F = 282.34, P < 0.0001)\). A post-hoc Tukey’s HSD test indicated a significant difference between the regression lines of the verteporfin and control groups \((P < 0.0001)\).

Analysis of the effects of YAP function inhibition on cyclic stretch proliferative response was also performed by culturing lenses at 6% stretch amplitude at 0.20 Hz with a verteporfin

Table 2. Labeling Indices of Lenses Cultured Under Varying Cyclic Strain Frequencies

| Strain Frequency, Hz | Labeling Index, % | n  |
|---------------------|------------------|----|
| 0                   | 3.23 ± 0.27      | 4  |
| 0.05                | 3.80 ± 0.51      | 4  |
| 0.10                | 5.52 ± 0.27      | 4  |
| 0.20                | 7.56 ± 0.23      | 4  |

Table 3. Labeling Indices of Verteporfin Treated and Paired Control Lenses Cultured Under Varying Static Strain Amplitudes

| Strain Amplitude (%) | Verteporfin | Control | n  |
|----------------------|-------------|---------|----|
| 0                    | 0.925 ± 0.24| 1.21 ± 0.10| 4  |
| 6                    | 1.27 ± 0.20| 3.86 ± 0.21| 4  |
| 12                   | 1.14 ± 0.14| 5.92 ± 0.59| 4  |
FIGURE 5. Variation in EdU labeling localization under different strain conditions. Qualitative analysis of representative images of lenses cultured under null strain (A), static strain (B), and cyclic strain (C–F), and stained for the EdU proliferative marker (green). The null (A) and static (B) stretch lenses are shown as mosaics going from one side of the equator to the other. The cyclic stretch lens (C–F) shows discrete images taken at different points along the anterior capsule: the anterior pole (C) and points along the equator (D–F). Lenses cultured under null strain (A) showed little reactivity with the EdU stain. In static strain (B), and cyclic strain (C) lenses, EdU labeling was primarily in the germinative zones (GZ). Magnified images of the GZ of null (A), static (B), and cyclic (F) stretched lenses are shown.
treated and untreated paired control (Fig. 7). The mean labeling index of the untreated control was 6.30 ± 1.58. The verteporfin treated group had a mean labeling index of 1.30 ± 0.33. A paired t-test showed a significant difference between the means (P < 0.0001).

The data shows that when YAP function was inhibited by verteporfin, the correlation between LEC labeling index and stretch amplitude was effectively eliminated. This suggests that YAP plays a crucial role in the transduction of mechanical signals into an upregulation of LEC proliferation. Furthermore, when YAP function is inhibited those signals are blocked and LEC proliferation does not increase with mechanical stretching.

**DISCUSSION**

This study was designed to determine if radially stretching the lens had an effect on LEC proliferation and whether YAP was involved in the mechanotransductive signaling pathway driving stretch-induced LEC proliferation. The stretch response was found to be dependent on both stretching amplitude and frequency; qualitative analysis of the localization of LEC proliferative activity showed differences between static and cyclic stretch. YAP was found to play an important role in the signaling pathway.
Localization was also not detected in the untreated (untreated lens (K) lenses cultured under null stretch conditions. Lenses were stained with the nuclear stain Hoechst (A, D, G, J) and for YAP (B, E, H, K). Nuclear localization of YAP was observed in the statically stretched untreated lens (C). This nuclear localization response was not observed in the statically stretched lens treated with verteporfin (F). YAP nuclear localization was also not detected in the untreated (I) or verteporfin treated (J) lenses cultured under null stretch conditions.

These results have important implications for understanding lens growth and morphogenesis, as well as approaches for modulating LEC proliferation. Controlling LEC proliferation could allow for retarding lens growth as a means for delaying presbyopia or cataract, as well as prevention of PCD (i.e., regeneration of the lens material following cataract surgery). Our data suggest that behavioral, environmental, and therapeutic approaches may be feasible for limiting or encouraging lens growth.

The human lens continues to grow throughout life, with an apparent bi-phasic growth pattern. It may be that the initial, very rapid, prenatal growth phase is driven by a rapid increase in lens capsule surface area and constant stretching forces, whereas the later, much slower, growth phase is retarded by the partial relief of lens stretching during accommodation. This is supported by Augusteyn's observation that the transition between growth phases occurs near the time of birth, as does the ability to accommodate. In non- or minimally accommodative species, age-matched lenses tend to be much larger (e.g., a 6-month-old pig lens may be ~400 mg, whereas an infant human lens is ~150 mg), possibly due to persistent disaccommodation.

Earlier studies have identified YAP as playing an important role in the regulation of tissue size, including the lens. When YAP is unphosphorylated and active, it is localized in the nucleus, acting as a transcriptional coactivator promoting the expression of genes inducing cell proliferation, survival, and migration. YAP is primarily regulated by the Hippo-signaling pathway, which when activated, phosphorylates YAP and inhibits its activity. The Hippo pathway is regulated by various mechanisms, including cell-cell contact, cell polarity, cellular energy status, hormonal signals and, most relevant to this study, mechanical cues. Studies on other cell types have observed that Hippo pathway activity is downregulated by stretching of those cells via the phosphorylation of the LATS1 kinase, which is the primary negative regulator of YAP. While the role of YAP in mechanosensing and cell proliferation has been studied in other cell types, and its expression in the lens is well documented, this study is the first to establish the link between mechanical cues and the regulation of the YAP protein in the lens.

The data presented in this study support the hypothesis that stretching the lens results in the activation of YAP and a subsequent increase in proliferative activity. However, YAP regulation is controlled by several different pathways with a significant amount of crosstalk between them and several other mechanosensing pathways may also be at play. In addition to the Hippo signaling pathway, RHO and MAPK/ERK signaling have also been implicated in the mechanoregulation of YAP. In other tissue types the signaling pathways Wnt, TGF-β, and Notch have been also been shown to increase cell proliferation in reaction to shear stress without the mediation of YAP. Additionally, p38 and JNK signaling pathways have been reported to respond to stretch and increase cell proliferation. Further studies will be necessary to determine what role the different mechanotransduction pathways play in the regulation of LEC growth.

This study demonstrates that stretching the porcine lens and connective tissues ex vivo results in increased LEC proliferation. There are several factors which could modulate LEC behavior during stretching. First, the capsule experiences increased tension due to the increase of zonular tension; this increase in capsule surface area will necessarily increase the footprint of LECs. LECs will also presumably experience increased apical pressure from the fiber cell bundle in the stretched state which could vary with position. LEC-LEC tensile forces may increase as well. Stretching could also drive fluid flow in and out of the lens or drive an increased rate of transport via convection due to relative motion of the capsule to the surrounding media. Finally, it is also possible that alternate signaling molecule(s) are activated or transported due to the stretching motion. Further investigation is required to pinpoint the underlying mechanism(s) of the stretch-induced change in proliferation.

The microscopy results presented in this study suggest increased labeling in cyclic and static stretch lenses compared to static lenses. The labeling indices measured using flow cytometry for the different stretch conditions were in qualitative agreement with microscopy findings. When visualizing the distribution of proliferative activity, the majority of EdU staining was observed at the GZ (Fig. 5). Future work will quantitatively assess spatial variations in proliferation.

Fluorescent microscopy showed activated nuclear YAP in LECs in the GZ (Fig. 8). This activation of YAP was only observed in stretched lenses uninhibited with verteporfin; YAP activation was observed in neither the unstretched lenses nor lenses treated with verteporfin. In some regions of the GZ of the statically stretched, EdU stained LEC proliferative activity (Figs. 5B, 5E) was observed in a similar density as the static stretched lens stained for YAP nuclear localization (Fig. 8B); suggesting a correlation between YAP nuclear localization and DNA synthesis. The microscopy technique used was unable to visualize cytoplasmic YAP and as a result provides no information on the total YAP content of the LECs. Additionally, the distribution of YAP nuclear localization likely changes depending on the position on lens capsule, and potentially corresponds to the local strain profile of the LECs. Further studies are needed to more completely characterize the behavior of YAP in response to stretching the lens.

While mechanotransduction pathways are highly conserved and every effort was made to replicate the microenvironment...
of the lens in vitro, caution should be used in extrapolating these findings to predict the in vivo behavior of human LECs. We kept the lens and connective tissues intact and retained the anterior vitreous attached to the lens. Still, the conditions experienced by the lens during the study differed from those in vivo. For example, cell culture media composition, including oxygen content, could alter the magnitude of the change in proliferation that was observed. Stretch amplitudes were chosen to match physiological extents of stretching: the maximal stretch-induced change in equatorial radius in a young human lens has been found to be between 5%–10%. In the present study, porcine lenses were used, which have different geometric and mechanical properties when compared to human lenses. Therefore, the distribution of mechanical stresses and the deformation from stretching likely differs between species. Additionally, pigs do not accommodate, resulting in a smaller and less robust ciliary muscle than that found in a human or primate eye. However, the biomolecular composition and crystallin distribution in both the human and porcine lenses are very similar. Further, mechanotransduction signaling and gene expression are highly conserved between species. Thus, if the LECs in the porcine mechanotransduction signaling and gene expression are highly conserved between species. Thus, if the LECs in the porcine lenses have an increased rate of proliferation in response to mechanical stretch, a similar response, albeit possibly with a different magnitude, would likely be observed in humans.

There are several possible explanations for the higher labeling index immediately after dissection relative to 24 hours of culturing after dissection. A stress response induced by changing the LEC microenvironment, subsequent acclimation of the LECs to the new environment, strain acting on the porcine lens in vivo and post mortem, and mechanical stimulation during the dissection process may contribute to the difference in labeling indices. Longer culture times could therefore be preferable for later studies in order to avoid any initial confounding cellular response to a new microenvironment or effects from mechanical loading prior to the culture period.

The results of the study establish a link between mechanical stretching and the upregulation of LEC proliferative activity, as well as identifying a target which, when inhibited, would reduce the growth of the lens. Overall this study provides new insights into the processes controlling lens growth and opens novel avenues by which to study the etiology of age-related vision disorders of the lens. Future work will map the localized proliferation changes with corresponding mechanical strains in the capsule.

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