Expression of Insulin-Like Growth Factor 2 Receptor in Corneal Keratocytes During Differentiation and in Response to Wound Healing

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Purpose. Insulin-like growth factor 2 receptor (IGF2R) associates with ligands that influence wound healing outcomes. However, the expression pattern of IGF2R and its role in the cornea is unknown.

Methods. Human keratocytes were isolated from donor corneas. Fibroblasts (fibroblast growth factor 2 [FGF2]-treated) or myofibroblasts (TGF-β1-treated) were analyzed for IGF2R and α-smooth muscle actin (α-SMA) expression by Western blotting and immunolocalization. Mouse corneas were wounded in vivo and porcine corneas ex vivo. The IGF2R and α-SMA protein expression were visualized and quantified by immunohistochemistry. The IGF2R gene expression in human corneal fibroblasts was knocked-down with targeted lentiviral shRNA.

Results. The IGF2R is expressed in epithelial and stromal cells of normal human, mouse, and porcine corneas. The IGF2R increases (11.2 ± 0.4-fold) in the epithelial and (11.7 ± 0.9-fold) stromal layers of in vivo wounded mouse corneas. Double-staining with α-SMA- and IGF2R-specific antibodies reveals that IGF2R protein expression is increased in stromal myofibroblasts in the wounded cornea relative to keratocytes in the normal cornea (11.2 ± 0.8-fold). Human primary stromal keratocytes incubated with FGF2 or TGF-β1 in vitro demonstrate increased expression (2.0 ± 0.4-fold) of IGF2R in myofibroblasts relative to fibroblasts. Conversion of IGF2R shRNA-lentiviral particle transduced corneal fibroblasts to myofibroblasts reveals a dependence on IGF2R expression, as only 40% ± 10% of cells transduced converted to myofibroblasts compared to 86% ± 3% in control cells.

Conclusions. The IGF2R protein expression is increased during corneal wound healing and IGF2R regulates human corneal fibroblasts to myofibroblast differentiation.

Keywords: cornea, wound healing, myofibroblasts, IGF2R, TGF-β1
IGF2R Expression in Corneal Wound Healing

IGF2R

uPAR & plasminogen

lysosomal enzyme-M6P

lysosomal enzyme-M6P

lysosomal enzyme-M6P

IGF2

TM domain

COOH

Figure 1. Schematic diagram of IGF2R protein. The IGF2R protein is a type I integral membrane protein with a single transmembrane (TM) domain. It contains 15 contiguous domains (gray and blue rectangles) in its extracellular region. Each domain is approximately 150 residues in length, except domain 13 is larger and contains a 48-residue fibronectin type II (FNII) insert (black). The domains are joined together by a short linker of 5 to 12 residues in length whereas the 15th domain is joined to the transmembrane region by a 25-residue linker. The locations of known ligand binding sites are shown (blue rectangles).

Lysosomal enzymes from the trans Golgi network to lysosomes; controlling extracellular levels of specific proteins through binding, internalization, and delivery of proteins from the extracellular milieu to lysosomes for degradation; and serving as a scaffold for proteins to interact at the plasma membrane surface. During development, the most important molecule that interacts with the receptor is IGF2, which binds to domain 11 and is delivered to the lysosome for degradation. By regulating the circulating levels of IGF2, a growth-promoting polypeptide that signals via the IGF1 and insulin receptors, IGF2 regulates the circulating levels of IGF2, a growth-promoting IGF2. In addition, knockdown (KD) of IGF2R in multiple cell phenotype that can be prevented by also knocking down Lysosomal enzymes and other proteins, such as renin and insulin, are developmentally regulated. Microarray analyses of the whole rat cornea 3 and 7 days after ex vivo plasminogen activator receptor (uPAR), plasminogen, connective tissue growth factor (CTGF), retinoic acid, and heparanase, through mechanisms yet to be elucidated. This receptor may serve as a scaffold for efficient activation of plasminogen by urokinase bound to uPAR through interaction of uPAR and plasminogen to domain 1 (Fig. 1). Binding of the receptor to a number of known molecules involved in corneal wound healing suggests IGF2R also is involved in this process.

The IGF2R is present in most tissues analyzed and its levels are developmentally regulated. No information is available concerning IGF2R protein expression with respect to the various cell types in the cornea. Furthermore, the ligand, IGF2, is present in aqueous humor and in stromal extracts, and IGF2 stimulates the proliferation of keratocytes in culture. Therefore, this study was performed to define the expression pattern of the IGF2R protein in the cornea and to determine whether IGF2R is required for corneal wound healing. The IGF2R was examined in vivo, ex vivo, and in vitro under normal conditions and in response to injury. In addition, a KD strategy was used to evaluate the role of IGF2R in corneal fibroblast differentiation to myofibroblasts.

Materials and Methods

Human Corneal Tissue and Cell Culture Conditions

Human corneas from deidentified organ donors were obtained from the Wisconsin Lions Eye Bank (Madison, WI, USA) within 48 hours of death. Our studies were conducted in compliance with the tenets of the Declaration of Helsinki. The use of deidentified tissue from nonliving individuals is not human subject research as described under section 45 CFR part 46 of the USA Code of Federal Regulations, and this exemption was recognized in writing by the Institutional Review Board.

All experiments were carried out using cells or tissue sections from at least four different donors. Human corneas for immunolocalization were fixed in formalin and embedded in paraffin. The epithelial and endothelial layers of other corneas were scraped from the stroma and the stromal cells were released by collagenase. Proteins were extracted from the cells of the three layers using lysis buffer (50 mM Tris, pH 7.4, 150 mM NaCl, 1% Triton X-100, 0.1% deoxycholate) containing protease inhibitors (Roche Diagnostics Corp., Indianapolis, IN, USA). Total protein was determined by the Bradford method. Primary human corneal epithelial cells were isolated from corneas using Dispase (25 caseinolytic units/ml; Life Technologies, Grand Island, NY, USA). Cells were cultured in defined keratinocyte serum-free medium (Life Technologies). Proteins were extracted from the cells as described above.

Human corneal stromal cells were cultured following removal of endothelial and epithelial layers from the donor corneas, and the stromal cells were released by collagenase digestion as described previously. The cultured fibroblast-like stromal cells were maintained in high glucose Dulbecco's modified Eagle's media (DMEM; Life Technologies) supplemented with 1% L-glutamine (Life Technologies) and 10% fetal bovine serum (FBS; Sigma-Aldrich Corp., St. Louis, MO, USA) at 37°C in a 5% CO2 atmosphere. Defined protease characteristic of fibroblasts and myofibroblasts were generated by seeding the stromal cells onto collagen (Advanced BioMatrix, San Diego, CA, USA)-coated wells in Defined medium (DMEM plus 1% RPMI vitamin mix; Life Technologies), 100 mM nonessential amino acids, 1 mM pyruvate, 100 μg/ml ascorbic acid, 1 mM glicine, 8 μg/ml insulin, 55 μg/ml transferrin, 5 μg/ml selenium, 100 μg/ml sodium selenite, and 10 μg/ml hydrocortisone. The cells were grown to confluence, and then serially passaged every 7-10 days. No information was available concerning IGF2R protein expression with respect to the various cell types in the cornea. Furthermore, the ligand, IGF2, is present in aqueous humor and in stromal extracts, and IGF2 stimulates the proliferation of keratocytes in culture. Therefore, this study was performed to define the expression pattern of the IGF2R protein in the cornea and to determine whether IGF2R is required for corneal wound healing. The IGF2R was examined in vivo, ex vivo, and in vitro under normal conditions and in response to injury. In addition, a KD strategy was used to evaluate the role of IGF2R in corneal fibroblast differentiation to myofibroblasts.

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acid), grown to confluence, and then treated for seven days by adding either 10 ng/mL fibroblast growth factor 2 (FGF2, promotes fibroblast phenotype; Life Technologies) or 1 ng/mL TGF-β1 (promotes myofibroblast phenotype; R & D Systems, Minneapolis, MN, USA) to the Defined medium.48

Murine Corneal Tissue

Our studies with mice adhered to the Association for Research in Vision and Ophthalmology (ARVO) Statement for the Use of Animals in Ophthalmic and Vision Research. This study was done in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals. The protocol was approved by the Institutional Animal Care and Use Committee (IACUC, AUA00002252), and buprenorphine was administered subcutaneously to minimize pain. The C57Bl/6J mice were anesthetized with isoflurane, and propa- racaine drops were instilled in the experimental eye. A trephine was used to outline a 2-mm circle on the cornea. To generate an epithelial scrape injury, the epithelial layer was removed within the outlined area using an Alger brush. After 7 days, the mice were euthanized, and the eyes were removed and fixed in 10% phosphate-buffered formalin.

Immunohistochemistry of Human and Murine Corneas

Formalin-fixed, paraffin-embedded human or murine corneal sections were deparaffinized, rehydrated, and treated with 10 mM citrate buffer containing 0.05% Tween 20 at pH 6 at 100°C for 10 minutes for antigen retrieval. After blocking, sections were incubated with IGF2R-specific antibodies (for human corneal sections, 1:200 dilution of product #PA3-850 polyclonal rabbit serum [ThermoFisher, Waltham, MA, USA] was used or for murine corneal sections, 1:200 dilution of B14.5 polyclonal rabbit serum23,39) or the corresponding preimmune serum at 4°C overnight. Biotinylated secondary antibody, Vectastain ABC-avidin alkaline phosphatase reagent, and Vector Red alkaline phosphatase fluorescent substrate (Vector Laboratories, Burlingame, CA, USA) were used for detection of the primary antibody. Sections were examined using a Nikon Eclipse 80i microscope equipped with a Nikon DS-FI1 camera and Elements D software (Nikon, Melville, NY, USA) for image capture. The Texas Red excitation and emission filters were used. Images were quantified using auto measure plus module of axiovision 4.8.1 software (Zeiss, Jena, Germany). The morphometric analysis was setup on a positive control slide using the pixel intensity of alkaline phosphatase stain to segment positively stained areas (thresholding); preimmune serum staining was used to set the threshold pixel intensity. The software also generates a postprocessed image of the segmented area for each image; thus, providing visual confirmation. For quantifying positive staining in either epithelial or stromal region, the data were corrected by manual removal method of the epithelial or stromal area. The percent stained area was calculated by measurement of the total alkaline phosphatase positive area above the threshold/total epithelium or stroma image area in square microns.

Porcine Corneal Organ Culture,
Immunohistochemistry, and Immunofluorescence Staining

Porcine eyes purchased from Pel-Freez (Rogers, AR, USA) were sterilized with iodine before performing a 5-mm anterior keratectomy. In this model, corneas are wounded by anterior keratectomy using a sharp-edged cylinder to create a wound through the epithelium, basement membrane, and the anterior stroma with subsequent removal of the incised tissue. Untreated or wounded corneas were removed with 2-mm scleral rims, mounted on an agar base containing 1% agarose and 1 mg/mL bovine collagen (Purcol; Advanced Biomatrix, Poway, CA, USA). The DMEM-F12 medium (Life Technologies) containing gentamycin and α-D-glucopyranosyl ascorbic acid was added to cover the sclera to the level of the limbus, and the air-exposed epithelium was moistened daily and the culture medium was changed every other day for 2 weeks as described.40 After 2 weeks, the corneas were fixed in formalin and paraffin embedded. Deparaffinized sections were incubated in 10 mM citrate buffer pH 6.4 at 100°C for 5 minutes for antigen retrieval. Following blocking for nonspecific binding, sections were prepared for either 5,5′-diaminobenzidine (DAB) and hematoxylin staining using horseradish peroxidase (HRP)- conjugated goat anti-rabbit antibody and DAB peroxidase substrate kit (Vector Laboratories) or immunofluorescence imaging (triple-labeled with antibodies specific for IGF2R (1:200 dilution of ThermoFisher product #PA3-850 and Alexa Fluor 488 Goat Anti-Rabbit IgG [Life Technologies] as the secondary antibody) and α-smooth muscle actin (α-SMA, Cy5 conjugated; Sigma-Aldrich Corp.), with nuclei stained with DAPI. Fluorescence microscopy was carried out using a Zeiss Axiosplan2 microscope at the Microscopy Shared Resource Facility at the Icahn School of Medicine at Mount Sinai with the filter sets ET 470/40x and ET525/50m for Alexa Fluor 488, and ET 620 and ET700/75m for Cy5. Using ImageJ (National Institutes of Health [NIH], Bethesda, MD, USA) threshold pixel intensity was set for all images, and the area in the stroma greater than the threshold measurement was quantified for each image. Values are reported as the mean ± SE.

Immunocytochemistry of Cultured Human Cells

Cultured human corneal fibroblasts and myofibroblasts were washed with PBS and fixed for 20 minutes on ice with 10% (vol/vol) formalin in PBS. After fixing, cells were incubated in PBS containing 0.1% Triton X-100 (vol/vol) for 5 minutes at room temperature. Fluorescein-conjugated antibody specific to α-SMA (Sigma-Aldrich Corp.) was diluted 1:100 in PBS containing 10 mg/mL BSA and 0.01 μg/mL Hoechst H33342 stain (Life Technologies) and incubated overnight at 4°C. The cells were washed with PBS and imaged on a Nikon Eclipse Ti fluorescent microscope equipped with a Nikon Digital Sight DS-Qi1MC camera using NIS Elements D3.10 ink software package. The LAMPI staining was done as above using a 1:200 dilution of the monoclonal antibody H4A3 developed by J. T. August and J. E. K. Hildreth (Developmental Studies Hybridoma Bank, University of Iowa, Iowa City, IA, USA, under the auspices of the National Institute of Child Health and Human Development, NIH) and a 1:1000 dilution of goat anti-mouse antibody conjugated to Alexa Fluor 568 (Life Technologies). The IGF2R was detected using IGF2R-specific antibody at a 1:100 dilution (ThermoFisher product #PA3-850) in PBS containing 0.1% BSA and 0.1% saponin and a 1:1000 dilution of chicken anti-rabbit antibody conjugated to Alexa Fluor 594 (Life Technologies). Double-staining for the detection of F-actin and α-SMA was conducted using a 1:200 dilution of fluorescein-conjugated phalloidin (Sigma-Aldrich Corp.) and a 1:200 dilution of α-SMA–specific antibody (Abcam, Cambridge, MA, USA) with a 1:500 dilution of goat anti-mouse antibody conjugated to Alexa Fluor 568 (Life Technologies), respectively. To compare expression levels between the fibroblast and myofibroblast phenotype for each of the above proteins, the same exposure time was used when imaging the stained cells. Negative controls without a primary antibody were included in each experiment as well as using the corresponding preimmune.
serum for the IGF2R-specific antibody. Cells were identified as myofibroblasts by analyzing approximately 200 to 500 cells per micrograph for the presence of assembled α-SMA fibrils.

**Western Blot Analyses**

The proteins from the dissected human corneal layers were extracted as described above and loaded onto SDS-PAGE gels in Laemmli SDS-PAGE sample buffer based on protein content. Cultured human stromal fibroblast and myofibroblast lysates were generated in Laemmli SDS-PAGE sample buffer and an equivalent number of cells (fibroblasts or myofibroblasts) were loaded into each lane of SDS-PAGE gels. The number of cells was determined by counting Hoechst H33342 (Life Technologies) stained nuclei using Image] software (version 1.46r; NIH). Nuclei were imaged using a Nikon Eclipse Ti fluorescent microscope and four random images were taken for each well. Proteins were transferred electrophoretically to Immobilon-P PVDF membrane (ThermoFisher) and subjected to quantitative Western blot analyses as described previously. Briefly, after incubating the membranes with antibodies specific to IGF2R (product #PA3-850; ThermoFisher), α-SMA (Abcam), or LAMP1 (Cell Signaling Technology, Inc., Danvers, MA, USA), the membranes were probed with horseradish peroxidase-conjugated Protein A (ThermoFisher) or goat anti-mouse IgG (ThermoFisher). The proteins were visualized using SuperSignal WestPico enhanced chemiluminescence reagents (ThermoFisher). The proteins were visualized using SuperSignal WestPico enhanced chemiluminescence reagents (ThermoFisher), and the resulting bands were quantified using a Bio-Rad ChemiDoc XR+ system and Image Lab software (Bio-Rad Laboratories, Inc., Hercules, CA, USA).

**Relative Quantitative Real-Time PCR**

Total RNA was extracted from cultured human corneal stromal fibroblasts and myofibroblasts derived from four human cornea donors using the RNaclyt kit (Qiagen, Valencia, CA, USA) and the amount of RNA isolated was quantified by optical density at 260 nm. RNA (1 μg) from each sample was reverse transcribed to cDNA using SuperScript III First-Strand Synthesis System (Life Technologies). Real-time PCR amplification of IGF2R cDNA was performed with previously published primers (forward primer, 5'-GAGGGAAGGCGAGAAGACG-3' and reverse primer, 5'-GTTCGCGGCTATCTGCAGC-3') by using Bullseye EvaGreen qPCR Master Mix (MidSci, St. Louis, MO, USA) with 6-carboxyl-X-Rhodamine (ROX) as the reference dye in an Applied Biosystems 7500 System (Life Technologies). The results of real-time PCR were analyzed by the comparative threshold cycle method and normalized to glyceraldehyde 3-phosphate dehydrogenase (GAPDH; forward primer, 5'-TCAAGCTATGGACGA-3' and reverse primer, 5'-ACCCGTGTCGATCGGACCT-3') as an internal control. The relative quantity of IGF2R was compared to the relative quantity of GAPDH for every sample. Each sample was run in triplicate. The DNA sequencing of each sample confirmed the identity of the amplified PCR product as IGF2R.

**Lentivirus-Mediated KD of IGF2R**

Knockdown of IGF2R expression was performed in primary human keratocytes using lentivirus to deliver short hairpin RNA interference (shRNAs). Lentiviral delivery of a shRNA expression cassette specific for the human IGF2R was performed using the sequence 5'-GCACCGGTATCGACCTT CTTAGAGAAAGATGATCTTGCCGG-3', a shRNA previously shown to efficiently reduce IGF2R protein expression in human cancer cells. The shRNA sequence was cloned into the AgeI and EcoRI sites of the Addgene plKO.1hygro plasmid 24150, which encodes a hygromycin resistance gene for selection (developed by R. Weinberg and obtained from Addgene, Cambridge, MA, USA) to make pLKOhygro6588. The insert was confirmed by DNA sequencing. Virus particles were produce by cotransfecting HEK293T cells with pLKOhygro6588 or pLKO.1hygro plasmid and helper plasmids using TransIT-293 transfection reagent (Mirus Bio LLC, Madison, WI, USA) according to the manufacturer’s protocol. Briefly, HEK293T cells were plated at 2 × 10^5 cells/well on 6-well plates for 24 hours before transfection. The pLKOhygro6588 or pLKO.1hygro plasmid and helper plasmids were incubated with the TransIT-293 reagent in Opti-MEM serum-free medium (Life Technologies) at room temperature for 30 minutes before addition to HEK293T cells. After 18 hours, the medium containing the transfection reagents was removed from the HEK293T cells and replaced with DMEM medium supplemented with 10% FBS and 1.1% BSA (viral collection medium). Following incubation for 24 hours, the medium was replaced with fresh viral collection medium and the cells were incubated for an additional 24 hours. The viral particles were concentrated from the medium by centrifugation at 17,500g for 3 hours at 4°C and resuspended in 0.1 times the original volume of DMEM, 10% FBS with 8 mg/mL polymyxin (Sigma-Aldrich Corp.) yielding a ×10 concentrated stock of viral particles. Transduction of human corneal fibroblasts to decrease IGF2R protein expression was conducted with freshly prepared viral particles. Briefly, primary corneal cells were plated at 4 × 10^5 cells/well on 6-well plates 24 hours before infection. Cells were washed with PBS and incubated with 1 mL of the concentrated viral particles. After 5 hours, 1 mL of DMEM medium was added and the cells were incubated for 18 hours. After 18 hours, the viral particle-containing medium was replaced with DMEM containing 10% FBS.

**Statistics**

Statistical analyses were performed using Sigma Plot 12.5 (Systat, San Jose, CA, USA). A 1-tailed Student’s t-test was performed for comparison of the differences in IGF2R immunostaining in wounded murine and porcine corneas versus nonwounded. This method also was used to compare the levels of mRNA for IGF2R in fibroblasts versus myofibroblasts. The protein levels of IGF2R, LAMP1, and α-SMA in cells treated with TGF-β1 versus FGF2 were compared to a hypothetical ratio of 1.0 using a 1-sample Student’s t-test. For comparison of the effects of knocking down IGF2R on the levels of IGF2R, LAMP1, and α-SMA, the wild type (WT) cell levels of the proteins was set to 100%, and the KD and pLKO vector control were calculated relative to the WT. The protein values for the IGF2R KD samples were compared to the pLKO vector control using the 1-tailed Student’s t-test. For comparison of the studies determining the effects of knocking down the IGF2R gene on conversion of corneal stromal cells to myofibroblasts, overall differences were determined by a 1-way ANOVA followed by multiple comparisons using the Holm-Sidak method. Data are presented as the mean ± SE.

**RESULTS**

**IGF2R Protein Expression in Corneal Tissue**

To evaluate the location of IGF2R within the normal cornea, immunohistochemistry was performed on tissue sections from human, mouse, and pig corneas. The results showed that IGF2R is present in the epithelial and stromal layers of normal, unwounded cornea (Figs. 2A, 3A, 4F). Most of the cells of the human epithelium stained for IGF2R; however, most of the staining for the mouse and porcine corneas was associated
with basal epithelium. In addition, Western blots of cultured primary human epithelial cells, and human corneal epithelial, stromal, and endothelial layer extracts confirmed the presence of IGF2R in all corneal layers (Figs. 2C, 2D).

We then asked if wounding altered IGF2R expression in the cornea in vivo. For these studies involving inflicted wounds we used a standard in vivo mouse epithelial wound model and a porcine organ culture model of corneal wounding, which removed the central epithelial layer and the anterior portion of the stroma. Mouse corneas were subjected to a 2-mm diameter epithelial scrape wound and allowed to heal for 7 days in vivo. Nonspecific preimmune staining of the wounded epithelial layer was much greater than observed for the nonwounded cornea. The IGF2R localization studies demonstrated that wounding the cornea increases IGF2R expression in the central epithelium by 11.2 ± 0.4-fold and stroma by 11.7 ± 0.9-fold (Figs. 3A, 3C). To confirm this result in another wounding model involving the stroma, the porcine corneal organ culture model was chosen because this model uses a large cornea that undergoes wound healing similar to that observed in the human.40 The porcine corneas were wounded by removal of a 5-mm trephined central portion of the epithelium, basement membrane, and stroma layers, and allowed to heal for 14 days in organ culture (Fig. 4). Consistent with the in vivo wounded murine cornea, a dramatic increase in IGF2R by 11.2 ± 0.8-fold was observed in the wounded porcine stroma compared to control (Figs. 4A, 4C, 4E).

Immunostaining using an Alexa Fluor 488–labeled secondary antibody for IGF2R was increased in the epithelium and stroma of the wounded cornea relative to that of the nonwounded cornea (Figs. 4F, 4G). This is consistent with the results in the murine model of corneal wounding (Figs. 3A, 3C). The epithelial staining for IGF2R was not observed using the colorimetric HRP-diaminobenzidine staining system (Figs. 4A, 4C), which is likely due to decreased sensitivity of this system relative to the Vector Red fluorophore used for the murine system (Figs. 3A, 3C) and the Alexa Fluor 488 fluorophore used for the porcine system (Figs. 4F, 4G).

We next asked if the IGF2R-positive cells in the cornea stromal layer were myofibroblasts. The differentiation of fibroblasts to myofibroblasts can be identified by the expression of α-SMA, and its assembly into stress fibers is an established marker of myofibroblasts in vitro and in vivo.42–44 Double-staining with α-SMA– and IGF2R-specific antibodies revealed that the dramatic increase in IGF2R expression observed in the stromal layer of the wounded porcine cornea occurs in myofibroblasts (Figs. 4F–K). An increase in the level of α-SMA also was noted in the wounded epithelial layer consistent with the increase in this protein during corneal epithelial-mesenchymal transition (EMT) following wounding.45,46
myofibroblasts, a process that can be recapitulated in stimulated at least in part by TGF-
During the wound repair process, corneal stromal fibroblasts
ALTERATION OF IGF2R EXPRESSION LEVELS BY TGF-
FIGURE 3. Immunodetection of IGF2R in tissue sections from wounded mouse cornea. Normal mouse corneas (A, B) or mouse corneas subjected to a 2-mm central scraped wound limited to the epithelial layer (C, D) were formalin-fixed, paraffin-embedded, and sections incubated with IGF2R-specific polyclonal antibody (A, C) or the corresponding preimmune serum (B, D). Biotinylated secondary antibody, avidin alkaline phosphatase reagent, and a fluorescent Vector Red alkaline phosphatase substrate were used for detection of the primary antibody. A representative experiment is shown of three independent replicates. Mouse corneas represent the unwounded central cornea (A, B) and the center of the wound bed (C, D). Scale bar: 40 μm. Arrows indicate representative cells in the stroma with intense staining for IGF2R. (E) Quantification of IGF2R staining area in the epithelial and stromal layers of control and wounded mouse samples is shown. Stained areas were above the threshold set based on the preimmune staining. The values represent the mean ± SE. *Indicates 1-tailed Student’s t-test, P < 0.005.

ALTERATION OF IGF2R EXPRESSION LEVELS BY TGF-β1–STIMULATED DIFFERENTIATION OF FIBROBLASTS TO MYOFIBROBLASTS IN CULTURE
During the wound repair process, corneal stromal fibroblasts
stated increased expression (2.0 ± 0.4-fold) of IGF2R in myofibroblasts relative to fibroblasts (Figs. 5A, 5B). These elevated IGF2R protein levels mirrored the 1.8 ± 0.2-fold increase in IGF2R transcript levels observed in myofibroblasts compared to fibroblasts (Fig. 5C). Immunofluorescence microscopy further supported the increased expression of IGF2R upon differentiation of fibroblasts to myofibroblasts (Fig. 5D). Taken together, these in vivo and in vitro results provided the first characterization of IGF2R protein expression in the cornea. Importantly, the results demonstrated upregulation of IGF2R protein during corneal wounding and in the process of fibroblast to myofibroblast differentiation. Furthermore, these results showed that alterations in IGF2R protein levels upon fibroblast to myofibroblast differentiation are mediated primarily by transcriptional regulation of the IGF2R gene.

Because one of the many cellular roles of IGF2R is delivering newly synthesized lysosomal enzymes to lysosomes, we asked whether the increased expression of IGF2R in myofibroblasts represents an expansion of endosomal/lysosomal compartments that may be needed for remodeling of the ECM by myofibroblasts. As an indirect measure of the number of endosomes/lysosomes, we quantified lysosome-associated membrane protein 1 (LAMP1), an established marker of late endosomes and lysosomes. The LAMP1 protein is among the most abundant lysosomal membrane proteins and its extensive glycosylation aids in protecting the luminal membrane of the lysosome from degradation by resident lysosomal hydrolytic enzymes. Western blot analysis demonstrated a 3.3 ± 0.9-fold higher amount of LAMP1 in myofibroblasts compared to fibroblasts (Figs. 5A, 5B). Immunofluorescence microscopy further supported the increased expression of LAMP1 upon differentiation of fibroblasts to myofibroblasts (Fig. 5D). The observed increase in LAMP1 levels is consistent with an expansion of the endosomal/lysosomal system in myofibroblasts compared to fibroblasts.

REQUIREMENT OF IGF2R FOR THE CONVERSION OF FIBROBLASTS TO MYOFIBROBLASTS
To begin to evaluate the role of IGF2R in the cornea, silencing of endogenous IGF2R in human primary corneal fibroblast-like cells was performed with lentiviral-based short hairpin RNA (shRNA) to reduce IGF2R transcript levels. Human corneal stromal cells can be transduced with lentivirus particles, and lentivirus delivery of shRNA is a method to silence gene expression by RNA interference (RNAi) in human corneal cells. Lentiviral delivery of an shRNA expression cassette specific for the human IGF2R was performed using an shRNA previously shown to reduce IGF2R protein expression in human cancer cells. The IGF2R KD experiments using human corneal stromal cells resulted in an approximately 50% reduction of IGF2R protein expression relative to cells transfected with an empty cassette (pLKO) and to WT cells (Fig. 6A).

In contrast to the parallel upregulation of IGF2R and LAMP1 upon TGF-β1–mediated conversion of fibroblasts to myofibroblasts (Figs. 5A, 5B), the levels of the lysosomal protein LAMP1 are not significantly different in the IGF2R KD cells when compared to control cells (Fig. 6A). These results showed that altering IGF2R protein levels do not affect the expression of LAMP1, indicating that TGF-β1–activated signaling pathways, but not IGF2R-mediated pathways, regulate the number of endosomal and/or lysosomal compartments.

We also evaluated the effect of decreased IGF2R levels on α-SMA expression by Western blot analysis. These data demonstrated approximately 25% less compared to cells transduced with the pLKO control vector and approximately 40% less α-
SMA in the IGF2R KD cells compared to WT control cells (Fig. 6A). Correspondingly, after 7 days in culture in the presence of TGF-β1, 86% ± 3% of cells transduced with the pLKO control vector lentiviral particles or 87% ± 3% of nontransduced (i.e., WT) cells converted to myofibroblasts as assessed by the presence of α-SMA–containing fibrils, whereas only 40% ± 10% of cells transduced with shRNA-IGF2R lentiviral particles were observed to express α-SMA–containing fibrils (Figs. 6B, 6C). Phalloidin staining shows that this decrease in α-SMA-containing fibrils is not due to a lack of F-actin fibrils in the IGF2R KD cells (Fig. 6B). Together, these results demonstrate IGF2R is required for corneal fibroblast upregulation of α-SMA synthesis, assembly of α-SMA–containing fibrils and differentiation to myofibroblasts.

DISCUSSION

The current report provides the first characterization of IGF2R protein expression in the cornea to our knowledge. The IGF2R protein is localized in corneal stromal and epithelial cells and its level is increased upon corneal injury. In the wounded cornea, a dramatic increase in IGF2R expression in the stromal layer occurs in myofibroblasts. The 2-fold higher IGF2R protein levels in myofibroblasts relative to fibroblasts in vitro correlates directly with the 1.8-fold increase observed in IGF2R transcript levels, indicating that alterations in IGF2R protein levels upon fibroblast to myofibroblast differentiation are due to transcriptional regulation of the IGF2R gene. A similar process occurs in the liver; upregulation of IGF2R transcription has been reported in hepatic stellate cells during fibrogenesis.51 Taken
together, the results demonstrated that elevation of IGF2R protein levels occurs as a consequence of corneal wounding and the process of TGF-β1–stimulated differentiation of fibroblasts to myofibroblasts.

Using a KD strategy, this study also shows that IGF2R is required for corneal fibroblast differentiation to myofibroblasts as assessed by assembly of α-SMA into fibrils, which is a characteristic feature of the myofibroblast phenotype. The IGF2R level correlates directly with the ability of α-SMA to assemble into fibrils as only 40% ± 10% of cells transduced with shRNA-IGF2R lentiviral particles converted to myofibroblasts compared to 86% ± 3% in cells transduced with the pLKO control vector lentiviral particles or 87% ± 3% in nontransduced cells. Similar findings were reported by Hao et al. using rat lung fibroblasts. In this study, differentiation of primary rat lung fibroblasts to myofibroblasts was induced by free silica, and the expression level of α-SMA and IGF2R was observed to be higher in myofibroblasts compared to fibroblasts.

These results lead to the question, “What are the possible mechanisms of action by which IGF2R could impact fibroblast to myofibroblast differentiation?” The IGF2R interacts with numerous M6P-containing and non-M6P–containing ligands that influence corneal biology and wound healing. In the in vivo mouse and ex vivo porcine models, IGF2R may be required for activation of latent TGF-β1 complex. This complex is composed of the TGF-β homodimer, the latency-associated peptide, and latent TGF-β–binding protein. One proposed mechanism for activation of this complex is through its binding to IGF2R via a M6P-containing N-glycan on the latency-associated peptide. Binding of the latent TGF-β complex to the receptor is thought to facilitate proteolysis by juxtaposing the cleavage site of the latency-associated peptide with IGF2R-bound proteases to release active TGF-β1.

The IGF2R protein must be critical for other mechanisms involved in TGF-β1–induced conversion of fibroblasts to myofibroblasts because in the cell culture studies reported here, active TGF-β1, which does not contain an M6P-
FIGURE 6. α-SMA fibril formation in primary human corneal myofibroblasts following transduction with control (pLKO) or shRNA-IGF2R lentiviral particles (KD). Lentiviral particle transduced or nontransduced cells (WT) were cultured in 1 ng/mL TGF-β1 for 7 days under serum-free conditions. (A) Lysates were generated and subjected to Western blot analysis (equivalent cell numbers loaded into each lane). Membranes were probed with IGF2R, α-SMA, or LAMP1-specific antibodies. Quantification is shown below the representative blots and is derived from corneal cultures obtained from four independent donors. Data are plotted as a percentage of the intensity of IGF2R, α-SMA, or LAMP1 bands in the shRNAi-IGF2R KD cells and cells transduced with pLKO control vector relative to nontransduced WT cells set to 100%. Values represent the mean ± SE. The Student’s t-test was
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used to compare the IGF2R KD cells to the pLKO control cells. *P < 0.02, n = 4. (B) Fixed cells were incubated with antibody specific to α-SMA that was visualized using a goat anti-mouse secondary antibody conjugated to Alexa Fluor 568 (red) and FITC-conjugated phalloidin (green). Nuclei were stained with Hoechst 33342 dye. An overlay of the phalloidin and α-SMA-stained cells is shown in the bottom. Representative micrographs are shown. Scale bar: 100 μm. (C) The images from the studies shown in (D) were quantified for the number of cells containing α-SMA fibrils. Approximately 200 to 500 cells per micrograph were scored for the presence of α-SMA-containing fibrils. The percentages shown are derived from four independent donors and are the average for the indicated number (n) of individual micrographs. The mean of the data derived from the four independent human donors also is shown. Values represent the mean ± SE. ANOVA, P < 0.001, individual comparisons use the Holm-Sidak method, *P = 0.002, ***P < 0.001.

containing glycan, was added to the fibroblasts, bypassing the activation step of latent TGF-β. One possible mechanism involves binding of uPAR to domain 1 of IGF2R; uPAR is a glycosylphosphatidylinositol (GPI)-anchored receptor composed of three domains and is involved in modulating extracellular proteolysis, cell adhesion, migration, and signaling.55,56 Bernstine et al.57 showed that conversion of corneal fibroblasts to myofibroblasts requires cleavage of the linker region of uPAR between domains 1 and 2, and reduced levels of surface bound uPAR.57 The IGF2R may facilitate uPAR cleavage by serving as a scaffold for orienting bound uPAR for cleavage by lysosomal proteases or plasmin also bound to IGF2R. Lysosomal proteases, such as cathepsin G, are known to cleave uPAR and to bind IGF2R through their M6P-containing carbohydrate complexes to domains 3, 5, and 9 of IGF2R.57–59 Plasminogen binds to domain 1 of IGF2R and can be activated to plasmin by upa-bound uPAR.56,57,60 Plasmin still bound to IGF2R can cleave and inactivate uPAR. Studies by Nykjaer et al.25 showed that uPAR is targeted by IGF2R to the lysosome for degradation. Thus, based on the requirement of uPAR cleavage for conversion of fibroblasts to myofibroblasts, the known functions of IGF2R and our data presented here, the mechanism for conversion of fibroblasts to myofibroblasts may require the following steps: IGF2R-facilitated uPAR cleavage, removal of cleaved uPAR from the cell surface by endocytosis of IGF2R-cleaved uPAR complex, trafficking of the complex to the lysosome, and degradation of uPAR. Future studies are needed to test this model.

The IGF2R also may be involved in CTGF-induced fibrosis. A recent study provided evidence that IGF2R is a receptor for CTGF.28 Because the effects of CTGF on abnormal extracellular matrix production are similar to TGF-β, CTGF is implicated in stimulating corneal scar formation.61 A lack of IGF2R may alter CTGF protein levels, thereby inhibiting myofibroblast formation. In addition, differentiation of rat kidney fibroblasts to myofibroblasts in culture has been shown to require a combination of TGF-β1, CTGF, and IGF2.62 Thus, IGF2R may have a critical role in orchestrating the appropriate levels of cytokines and growth factors required for corneal homeostasis and the conversion of fibroblasts to myofibroblasts during wound healing.

The data presented here would suggest inhibition of the synthesis of IGF2R would decrease conversion of fibroblasts to myofibroblasts and could be used to control corneal fibrosis; however, this may adversely alter corneal wound healing because of other functions of the receptor. One of the major roles of the receptor is to bind and transport proteins to the lysosome. This property would be important for the control of extracellular proteins that bind IGF2R and are involved in wound healing: IGF2 which regulates cell proliferation,55 M6P-containing small leucine-rich glycoproteins,55 including lumican, which is involved in inflammatory and wound healing processes as well as maintaining corneal transparency,56,57 and lysosomal enzymes, which if abnormally secreted degrade extracellular proteins. Because of these important functions of IGF2R, a better approach to controlling myofibroblast formation may be the design and use of molecules that specifically target the interaction of IGF2R with ligands involved in the conversion of fibroblasts to myofibroblasts and in the fibrotic process. For example, small molecules could be identified and used to specifically target the uPAR-binding site in domain 1 (Fig. 1), leaving other critical functions of this receptor intact. Given the multifunctional nature of IGF2R, future studies are warranted to investigate the mechanisms by which IGF2R can influence corneal biology and wound healing.

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