Inhibition of Human Corneal Myofibroblast Formation

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Corneal fibrosis is basically the wound-healing response of the cornea to different types of injuries, which fails to properly repair and preserve corneal structure. Corneal wounds, which result from trauma or refractive surgery, trigger a complex healing process that involves inflammation, activation of cell-signaling events, generation and release of numerous cytokines, apoptosis, proliferation, migration, adhe-
sion, differentiation, and remodeling of the extracellular matrix (ECM). Some of the obvious changes in the corneal stroma during this process are as follows: (1) keratocyte death in the wound area; (2) quiescent keratocyte activation to fibroblasts in the area adjacent to the wound site; (3) proliferation of fibroblasts and replenishment of the wound bed to remodel the wounded ECM; and (4) fibroblast, or other cell type, transformation to myofibroblasts, which are a contractile stromal cell phenotype often identified by the presence of α-smooth muscle actin (αSMA) fibrils contained within the cell. The formation of myofibroblasts and the increased deposition of fibrotic proteins in the ECM, allow for the cornea to rapidly repair the wound gap; however, this wound-healing process results in a less than optimal tissue, namely an opaque cornea that interferes with vision that may even cause blindness. The current solution to restore vision for most of these eyes is through corneal transplant surgery, which replaces the dysfunctional cornea with a healthy donor cornea. However, the global shortage of transplant-grade donor corneal tissue greatly restricts the number of corneal transplantations performed. Therefore, there is enormous clinical benefit to studying, understanding the mechanisms involved, and finding ways to prevent scar formation in corneal wound healing. One method of preventing fibrosis may be by controlling the key step(s) in wound healing, such as blocking the transforming growth factor-beta (TGF-β)-signaling pathway with the p38 inhibitor (p38inh), a highly selective, potent, and cell-permeable compound that inhibits p38 MAP kinase (p38MAPK). If successful, this potentially would alleviate the increasing demand for transplantable corneal tissue and treat other corneal injuries, which in the United States alone are estimated to be over one million annually. Of these various growth factors, TGF-β1 (T1), one of three TGF-β isoforms, has a central role in severe fibrotic diseases. T1 has been found to stimulate myofibroblast transformation in

METHODS. In HCF, Trx-SARA (Smad-pathway inhibitor) was used to block the TGF-β/Smad-signaling pathway, and the p38 inhibitor (p38inh, SB202190) was used to inhibit p38MAPK, thus blocking the TGF-β/p38-signaling pathway. HCF ± Trx-SARA or Trx-GA (SARA control) were serum starved overnight in Eagle’s minimum essential medium (EMEM) ± p38inh, grown in EMEM ± T1 ± p38inh for 24 hours, and then processed for indirect-immunofluorescence, Western blot, or quantitative real-time polymerase chain reaction to examine α-smooth muscle actin (αSMA) and other fibrotic genes, such as fibronectin, thrombospondin1, and type III collagen. In addition, the morphology and the effect of p38inh on myofibroblast phenotype after myofibroblast formation were examined.

RESULTS. We observed that Trx-SARA had little effect on αSMA expression, indicating that blocking the Smad pathway did not significantly inhibit myofibroblast formation. However, p38inh did significantly inhibit αSMA and other fibrotic genes, thus efficiently preventing the transition of HCFs to myofibroblasts. In addition, morphology changed and αSMA decreased in myofibroblasts exposed to p38inh medium, as compared with controls.

CONCLUSIONS. HCF transition to myofibroblasts was mainly through the p38 pathway. Therefore, blocking the p38 pathway may be a potential therapeutic tool for human corneal fibrosis prevention/treatment, because it controls myofibroblast formation in human corneal cells, while leaving other functions of T1 unaffected.

Keywords: fibrosis, myofibroblasts, corneal fibroblasts, TGF-beta signaling pathway, p38 pathway
comprised of the rigid scaffold Trx (the Escherichia coli essential medium [EMEM; ATCC, Manassas, VA, USA] plus 10% placed in 6-well plates in regular medium (RM; Eagle’s minimum from review by the Institutional Human Studies Committees at The study’s experimental protocols were judged to be exempt Declaration of Helsinki. Human corneas were obtained from the methods used in these studies adhered to the tenets of the human corneal stromal cells. By doing so, we hoped to find a to dissect the signaling response of fibrotic genes to T1 in cornea, while leaving other functions of T1 unaffected. signalling pathway. In our previous studies, we found that different T1-target proteins, even the same target protein as well as ECM synthesis and degradation. Also, T1 exhibits proliferation, adhesion, and migration in a variety of cell types, growth factor that has pronounced effects on cell growth, different pathways. It is known that T1 is a multifunctional in different human corneal cell types, were regulated through that different T1-target proteins, even the same target protein in vivo and in vitro, and by binding to its receptors on the cell surface, T1 triggers signaling, which leads to the activation of either a non-Smad-signaling pathway, such as p38(MAPK), Rab5(MAPK), PP2A, RhoA, and JNK, or the better-known Smad-signaling pathway. In our previous studies, we found that TGF-β signaling was involved in corneal wound repair in debridement, keratectomy, and penetrating wound models, and the type of wound or extent of injury affected which TGF-β-signaling pathway was stimulated. In addition, we found that different T1-target proteins, even the same target protein in different human corneal cell types, were regulated through different pathways. It is known that T1 is a multifunctional growth factor that has pronounced effects on cell growth, proliferation, adhesion, and migration in a variety of cell types, as well as ECM synthesis and degradation. Also, T1 exhibits both exacerbating and ameliorating features, depending on the phase of disease and site of action. The aim of this study was to dissect the signaling response of fibrotic genes to T1 in human corneal stromal cells. By doing so, we hoped to find a way to prevent the stromal cell transition to myofibroblasts by blocking specific TGF-β-signaling pathway(s) in the human cornea, while leaving other functions of T1 unaffected.

**Materials and Methods**

**Human Corneal Fibroblast Isolation**

Human corneal fibroblasts (HCF) were isolated from human corneal stromas, as described previously. All procedures/methods used in these studies adhered to the tenets of the Declaration of Helsinki. Human corneas were obtained from the National Disease Research Interchange (Philadelphia, PA, USA). The study’s experimental protocols were judged to be exempt from review by the Institutional Human Studies Committees at the Schepens Eye Research Institute/Massachusetts Eye and Ear. Briefly, human corneal stromal explants from donor eyes were placed in 6-well plates in regular medium (RM; Eagle’s minimum essential medium [EMEM; ATCC, Manassas, VA, USA] plus 10% fetal bovine serum [Atlanta Biologicals, Flowery Branch, GA, USA]), and incubated at 37°C with 5% CO2 until sufficient amounts of HCF migrated from the explants.

**Production of HCF Cell Lines**

We produced two cell lines (Trx-SARA-HCF and Trx-GA-HCF) by infecting HCF with a retrovirus containing either a Trx-SARA or Trx-GA plasmid, as described previously. Trx-SARA is comprised of the rigid scaffold Trx (the Escherichia coli thioeductaxin A protein) followed by the Smad-binding domain of SARA, which is a constrained 56-amino acid Smad-binding motif from the SARA (Smad anchor for receptor activation) protein. Trx-SARA blocks the Smad-signaling pathway by binding to the monomeric Smad proteins, thus reducing the level of Smad2 and 3 in complex with Smad1, after TGF-β stimulation. Trx-GA is a control Trx aptamer of Trx-SARA, which contains an 11-amino acid repeat of Gly-Ala.

**Table 1. Three Additional p38 Inhibitors Tested**

| p38 Inhibitor | Conc. | IC50* | References |
|---------------|-------|-------|------------|
| SB202190      | 10 μM | p38α = 280 nM | Manufacturer’s Data Sheet*4 |
| Birb796       | 10 μM | p38α = 38 nM | Gicenas J. et al. 2017*44 |
| SB239063      | 20 μM | p38α = 44 nM | Barone FC. et al. 2001*55 |
| SB205580      | 5 μM  | p38αβ = 0.6 μM | Bain J. et al. 2000*56 |

* Concentration, concentration.

*Inhibitory concentration, concentration of an inhibitor where the response is reduced by half.

**Cell Culture**

HCF ± Trx-SARA or Trx-GA were seeded in 100-mm dishes or chamber slides in RM until they reached 60% to 70% confluency, at which time, the cells were cultured either in basic medium (BM; EMEM only) to serum starve the cells overnight or in p38inh (SB202190; Sigma-Aldrich Corp., St. Louis, MO, USA) medium (BM + 10 μM p38inh) to serum starve and pretreat the cells with p38inh overnight. The concentration of p38inh SB202190 was determined based on published reports. The next day, cells were treated in BM ± 2 ng/mL T1 (R&D Systems, Inc., Minneapolis, MN, USA) ± 10 μM p38inh for 24 hours. Cells were examined and brightfield images were obtained (Carl Zeiss Microscopy, LLC, Thornwood, NY, USA). Then, they were either harvested by trypsinization for Western blot (WB) or quantitative real-time polymerase chain reaction (qRT-PCR) or fixed in cold methanol for indirect immunofluorescence (IF). Experiments were performed in at least triplicate for each condition. To confirm the specificity of p38inh (SB202190), three additional p38inhs (Table 1) were randomly selected and tested in the same manner. All p38inhs were found to have similar results.

In addition, HCFs were cultured in RM until they were 70% confluent, at which time they were stimulated to become myofibroblasts by exposing them to RM ± 2 ng/mL T1 for 3 days. The T1 cells were then rinsed with PBS three times, split into four groups, and cultured in the following for 1, 2, 3, or 4 days: (1) RM, (2) RM + 2 ng/mL T1, (3) RM + 2 ng/mL T1 + 10 μM p38inh, (4) RM + 2 ng/mL T1 + 10 μM p38inh. Media was changed every day and any media that included T1 had freshly prepared T1 added every day to ensure similar T1 activity throughout the experiment. Cells in different culture conditions were collected at each time point.

**SDS-PAGE and WB Analysis**

SDS-PAGE and WB analysis were performed as previously described. In brief, protein from cells was extracted with radioimmunoprecipitation assay buffer (10 mM Tris, 150 nM NaCl, 1% deoxycholic acid, 1% Triton X-100, 0.1% SDS, 1 mM EDTA; Sigma-Aldrich Corp.) plus protease inhibitors (aprotinin, phenylmethylsulfonyl fluoride, and sodium orthovanadate; Sigma-Aldrich Corp.). Protein samples of equal volume and protein were loaded onto gradient gels (8%–16% Tris-glycine gels; Invitrogen, Carlsbad, CA, USA), transferred onto nitrocellulose membrane (Bio-Rad; Hercules, CA), and incubated with primary antibodies (SmA (Abcam, Cambridge, MA, USA), p38α, phosho (p)-p38α (Santa Cruz Biotechnology, Inc., Dallas, TX, USA), cellular fibronectin (cFN; Sigma-Aldrich Corp.), thrombospondin-1 (TSP1; Thermo Scientific, Waltham, MA, USA), or β-actin (Sigma-Aldrich Corp.). Protein bands were viewed on an infrared imaging system (Odyssey; Li-Cor, Lincoln, NE, USA) by detecting secondary antibodies conjugated with IRDye 680RD or 800CW (Li-Cor). β-actin was used as an internal control.

**Indirect Immunofluorescence**

Cells were fixed with methanol and processed for IF, as described previously. In brief, cells were blocked (1% bovine

References

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serum albumin + 0.1% Triton X-100 (Sigma-Aldrich Corp.), incubated overnight at 4°C with anti-aSMA (Dako North America, Carpinteria, CA, USA), and then incubated for 1 hour at room temperature with the appropriate secondary antibody conjugated to fluorescein (Jackson ImmunoResearch, West Grove, PA, USA). Samples were mounted with mounting media containing 4’,6-diamidino-2-phenylindole (Vectashield; Vector Labs, Burlingame, CA, USA), a nuclear counterstain, and observed and photographed (Nikon Eclipse E800 equipped with an Andor Clara E camera and Nikon NIS Elements for Basic Research; Micro Video Instruments, Avon, MA, USA). Negative controls, where the primary antibody was omitted, were run with all experiments.

Quantitative Real-Time Polymerase Chain Reaction

As previously described,39 qRT-PCR was performed to quantify the fibrosis-related genes aSMA and type III collagen (Col III). In brief, total RNA was extracted from HCF (Trizol; Thermo Fisher Scientific, Waltham, MA, USA), purified (Qiagen RNeasy Mini isolation kit; Qiagen, Inc.; Valencia, CA, USA), and reverse transcribed to cDNA (High Capacity cDNA Reverse Transcription Kit; Applied Biosystems, Carlsbad, CA, USA). The qRT-PCR SYBR Green assay was performed with cDNA in master mix (KAPA SYBR Fast qPCR master mix; KAPA Biosystems, Wilmington, MA, USA) with primers for aSMA, Col III, and β-actin (Table 2; CCIB DNA Core Facility at Massachusetts General Hospital, Cambridge, MA, USA). Amplification was performed (Eppendorf multiplex 2 real time PCR machine; Eppendorf, Hauppauge, NY, USA) with the following thermal cycling conditions: 2 minutes at 50°C, 10 minutes at 95°C, 40 cycles of 15 seconds at 95°C, and 1 minute at 60°C. Relative mRNA levels of specific genes were reported as 2(−ΔΔCt).

TABLE 2. qRT-PCR Primer List and Sequences

| Gene       | Forward Primer (5′⋯3′)                     | Reverse Primer (5′⋯3′)                |
|------------|-------------------------------------------|--------------------------------------|
| aSMA       | GGTGGACGAAGCAGAGCAA                       | CAGTGGTGATGGCCATGTGTT                |
| Col III    | GGTGCCTGGGTTAATACGC                       | TCCAGGGATCCGGCAAGTT                  |
| β-actin*   | GACACTGTTGCTGGGTGACG                      | AGCACGTGTGTTGGGAGTG                  |

* VHPS-110 (Real Time Primers; Elkins Park, PA, USA).

RESULTS

aSMA- and TGF-β-Signaling Pathways

In initial studies, we used a stable Trx-SARA-expressing HCF cell line, Trx-SARA-HCF, to examine the role of the Smad pathway on aSMA expression in HCF. As seen in Figure 1, little, if any, aSMA expression was detected in all three HCF samples (HCF, TRX-GA, and Trx-SARA) when cultured in BM only; however, with the addition of T1, aSMA expression increased significantly in all three samples (**P < 0.01). Interestingly, the existence of Trx-SARA had little, if any, effect on aSMA expression, indicating that blocking the Smad pathway did not significantly inhibit aSMA protein expression in HCF. This suggests that T1 does not stimulate corneal fibrosis through the Smad pathway, but by some other means.

We then examined the role that the TGF-β/p38-signaling pathway plays on the expression of aSMA in HCF. First, we needed to confirm that the p38inh SB202190 could block the p38 pathway. As seen in Figure 2, we confirmed by WB the inhibitive effect of SB202190 on the phosphorylation of p38 in HCF and we calculated and graphed (Fig. 2) the ratio of phospho (p)-p38/p38 in HCF treated with the following: (1) BM, (2) BM + T1, and (3) BM + T1 + p38inh. Compared with BM samples (Fig. 2, inset lanes 1–3), the phosphorylation of β-actin was used as the loading control.

**Figure 1.** Analysis of TRX-SARA’s effect on aSMA expression in HCF. HCF ± Trx-SARA or Trx-GA were either treated with EMEM (BM) or BM + 2 ng/mL of T1, and then examined for aSMA protein expression by WB. In all samples, aSMA protein expression increased significantly with T1 stimulation (**P < 0.01). Insert is a representative blot from three independent experiments. β-actin was used as the loading control.
p38 in the T1 samples (Fig. 2, inset lanes 4–6) was significantly upregulated (\( **P < 0.01 \)). However, the addition of p38inh to HCF treated with T1 (Fig. 2, inset lanes 7–9), effectively blocked the phosphorylation of the p38 protein by endogenous T1 to levels that were undetectable by WB (data not shown). These data indicate that p38inh effectively blocks the TGF-\( \beta /p38 \) pathway in HCF.

Since T1-treated HCF transition to myofibroblasts, we observed if p38inh altered the cellular morphology and \( \alpha \)-SMA localization in HCF by treating them as follows for 24 hours: (A) BM only, (B) BM + T1, (C) BM + p38inh, and (D) BM + T1 + p38inh. As seen in Figure 3, the cells appeared to be spindle shaped in both BM control (Fig. 3A) and BM + p38inh (Fig. 3C), similar to what would be expected for HCF when T1 was added to BM (Fig. 3B), the cells changed from the fibroblastic spindle shape to a more myofibroblastic morphology, with stretched cellular hypertrophy and typical stress fibers.43,44 Surprisingly, we found that the morphology of HCF in BM + T1 + p38inh (Fig. 3D) was similar to BM control and BM + p38inh (Figs. 3A, 3C). Therefore, the presence of p38inh in the T1 samples appeared to efficiently inhibit the morphologic transition of HCF to myofibroblasts by exogenous T1 (Fig. 3D).

The \( \alpha \)-SMA localization and protein expression shown in Figure 4 agrees with the morphologic data (Fig. 3). As with Figure 3, BM control and BM + p38inh had only a few \( \alpha \)-SMA-labeled cells (Fig. 4A.a, 4A.c) and little \( \alpha \)-SMA protein expression (Fig. 4B, lanes 1 and 3); however, with T1 treatment (Figs. 4A.b, 4B, lane 2), there were more \( \alpha \)-SMA-labeled cells and a significant increase in protein expression (\( **P < 0.001 \)). By treating the HCF with BM + T1 + p38inh (Fig. 3D) was similar to BM control and BM + p38inh (Figs. 3A, 3C). Therefore, the presence of p38inh in the T1 samples appeared to efficiently inhibit the morphologic transition of HCF to myofibroblasts by exogenous T1 (Fig. 3D).

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Expression of Other Fibrosis-Related Genes and the p38 Pathway

The expression of certain fibrosis-related genes, such as cFN, TSP1, and Col III, has been shown to be upregulated in myofibroblasts at the mRNA and/or protein level.37,45,46 In addition, TSP1 protein levels have been shown to significantly decrease when exposed to p38inh.28 To examine if cFN and Col III were stimulated by T1 through the p38 pathway and when the p38 pathway is inhibited, quantitative RT-PCR (qRT-PCR) was performed. As with \( \alpha \)-SMA, cFN protein and Col III mRNA expression significantly increased in T1-treated HCF (Figs. 7A, 7B), \( *P < 0.05 \), as compared with control (BM only). When HCFs were treated with BM + T1 + p38inh, cFN protein (Fig. 7A, lane 4) and Col III mRNA expression (Fig. 7B) were greatly reduced, as compared with BM + T1 (\( *P < 0.05 \)). In fact, cFN protein and Col III mRNA expression returned to control levels (BM only) or lower. Interestingly, unlike \( \alpha \)-SMA, cFN, and Col III, TSP1 did not return to control levels or lower, but rather only decreased by approximately one-third, indicating that another pathway may also be involved with activating TSP1.28 These data indicate that cFN, TSP1, and Col III expression stimulated by T1 in HCF were at least partly regulated through the p38 pathway.

Role of p38 Inhibitor on Human Corneal Myofibroblast Phenotype

Since p38inh appeared to successfully inhibit myofibroblast formation due to T1 stimulation of HCF, we examined if the p38 inhibitor could not only reduce, but also reverse \( \alpha \)-SMA levels, thus changing the myofibroblast phenotype. As seen in Figure 8 (inset), the WB results showed that \( \alpha \)-SMA levels in T1-treated HCF (Fig. 8A, lane 2) were significantly higher than in the HCF control (\( P < 0.001 \); Fig. 8A, lane 1). On day 1 (D1), \( \alpha \)-SMA levels in both RM and p38inh groups continued to increase (Fig. 8A, lanes 3 and 4), compared with the T1-control (Fig. 8A, lane 2). On day 2 (D2), the \( \alpha \)-SMA levels in the RM sample leveled out (Fig. 8A, lane 5); however, in the p38inh sample (Fig. 8A, lane 6), the \( \alpha \)-SMA levels decreased and were
significantly lower than the RM sample at D2 \((P < 0.05)\). On day 3 \((D3)\), \(\alpha\)SMA levels in both media samples decreased (Fig. 8A, lane 7 and 8). Although both sample groups decreased, the \(\alpha\)SMA level in the p38inh sample continued to be lower than that in the RM sample, which returned to T1 level. As seen in Figure 8B, the mRNA data agreed with the protein data (Fig. 8A), in that T1-treated HCF mRNA expression was higher than the HCF control, and when the T1 samples were switched to RM only, the \(\alpha\)SMA mRNA expression continued to increase at D1 and D2, appeared to peak at D2, and returned to almost T1 levels by D3. Interestingly, the \(\alpha\)SMA mRNA expression of the samples that were switched to p38inh hovered around T1 levels at D1 and D2 and then decreased to control HCF \(\alpha\)SMA mRNA levels by D3. These data indicate that although the human corneal myofibroblasts in RM returned to T1-control levels 3 days after the removal of T1 from the culture medium, the presence of the p38 inhibitor accelerated the change of the myofibroblast phenotype (Fig. 8B.D2).

In Figure 9, HCF samples where T1 was continuously present in the medium were observed. As seen in Figure 9A.a–k, the morphology of HCF cultured in RM only (Figs. 9A.c, 9A.f, 9A.i) for D1 to D4 remained relatively similar to HCF control (Fig. 9A.a); however, the T1-treated HCF became increasingly more myofibroblastic with time (Fig. 9A.d, 9A.g, 9A.j). By D4 with p38inh (Fig. 9A.k), the morphology of the HCF appeared to revert to that seen in the HCF control (Fig. 9A.a). Localization of \(\alpha\)SMA was also examined at D4 (Fig. 9A.l–n) and the results agreed with the morphologic data (Fig. 9A.a–k). The D4 HCF control (Fig. 9A.l) had little to no \(\alpha\)SMA localization, and T1 samples (Fig. 9A.m) had an increase in \(\alpha\)SMA-positive cells. Interestingly, \(\alpha\)SMA localization in T1 + p38inh samples (Fig. 9A.n) was similar to that of HCF controls (Fig. 9A.a). No staining was observed in negative controls (data not shown). These samples were also examined for \(\alpha\)SMA protein levels (Fig. 9B), which increased when T1 was added to the medium. This increase continued to rise over time (T1-D3 compared with T1-control, \(P < 0.001\)); however, in samples with T1 + p38inh medium, \(\alpha\)SMA levels increased on D1, decreased by D2, and significantly decreased by D3, as compared with T1-control \((P < 0.05)\) and T1-D3 \((P < 0.01)\). These data indicate that blocking the p38 pathway changed the phenotype of human corneal myofibroblasts.

**DISCUSSION**

It is hard to completely avoid corneal injury, since injury can be caused by disease, accident, and even incisional surgery. Improper healing of these corneal wounds can lead to visual interference or even loss. In normal corneas, keratocytes are quiescent and structurally flattened cells found in the stroma; however, in wounded corneas, these cells are activated to become either fibroblasts or myofibroblasts, with the latter causing corneal fibrosis. T1, a key multifunctional growth factor that stimulates stromal cell transformation, is upregulated in corneal cells after a corneal wound and released into the stroma.\(^{47,48}\) The T1 then binds to its receptors on the cell surface and produces a cascade effect via various intracellular signaling pathways, which activate the keratocytes to phenotypically change, ultimately transforming into myofibroblasts with high levels of \(\alpha\)SMA. In addition to being associated with the development of stress fibers (\(\alpha\)SMA) during corneal repair, T1 is also associated with cell proliferation and migration. Indeed, our previous studies have shown that T1 proliferation can be greatly inhibited by blocking the Smad pathway.\(^{28}\) Therefore, to control the transition of HCF to myofibroblasts during wound repair while leaving the other functions of T1...
unaffected is extremely important and, we hope, would improve corneal healing results. To discern which pathway controls myofibroblast transition, two types of inhibitors were used in our studies to block either the TGF-β/Smad (Trx-SARA)- or TGF-β/p38 (p38inh, SB202190)-signaling pathways. After T1 stimulation, αSMA localization, as well as protein and mRNA expression, increased; however, no obvious change in αSMA expression, as compared with controls, was observed in HCFs containing Trx-SARA (Fig. 1), indicating that T1 did not affect the transition of HCFs to myofibroblasts through the Smad pathway. Surprisingly, when T1-stimulated HCFs were also treated with p38inh, not only did the morphology remain similar to controls (Fig. 3), but also the αSMA localization (Fig. 4A), protein expression (Fig. 4B), and mRNA expression (Fig. 5). This indicates that αSMA expression is regulated by T1 in HCF through the p38 pathway and by blocking the p38 pathway, the transition of HCF to myofibroblasts can be prevented, while leaving other T1 functions that occur through other signaling pathways (i.e., proliferation) unaffected.

The cornea is a transparent tissue, and the balance between ECM degradation and deposition in the stroma must be tightly regulated to maintain corneal function. Myofibroblasts are responsible for fibrotic-associated protein production, as well as ECM deposition and organization in wound healing.37,45,46,49 If we are able to diminish this fibrotic-associated protein expression by preventing stromal cells in a

**Figure 4.** IF and WB analysis of the p38 inhibitor’s effect on αSMA expression in HCF (A) A set of representative IF images of HCF treated in the following medium: (a) BM, (b) BM + T1 (2 ng/mL), (c) BM + p38inh (10 μM), (d) BM + T1 + p38inh. Little, if any, αSMA localization was observed in controls: BM + p38inh (a, c). As expected, αSMA localization increased with T1 stimulation (b); however, the presence of p38inh efficiently inhibited T1’s role in αSMA protein expression (d). Red, αSMA; Blue, 4’,6-diamidino-2-phenylindole, a nuclear counterstain. Scale bar: 100 μm. (B) αSMA protein expression increased significantly in HCF with T1 stimulation (lane 2; ***P < 0.001), as compared with controls: BM + p38inh (lanes 1 and 3); however, the addition of p38inh significantly inhibited T1’s role on the αSMA protein levels (lane 4; **P < 0.001), returning them back to control levels. Representative blot from one of three independent experiments is shown. β-actin was used as the loading control.

**Figure 5.** qRT-PCR analysis of p38 inhibitor’s effect on αSMA expression in HCF. qRT-PCR was performed on HCF that were treated in the following medium: BM, BM + T1 (2 ng/mL), BM + p38inh (10 μM), and BM + T1 + p38inh. The relative mRNA expression of αSMA in each group was corrected with β-actin, an internal control. The role of exogenous T1 on αSMA mRNA expression in HCF was significantly prevented with the addition of p38inh (*P < 0.05).

**Figure 6.** WB analysis of three additional p38 inhibitors’ effect on αSMA expression in HCF. The αSMA protein expression in HCF cultured with EMEM ± p38 inhibitor (p38inhns: 063 [SB239063]; Birb [Birb796], or 580 [SB203580]), or EMEM + 2 ng/mL T1 ± p38inh was analyzed. With the addition of T1 to samples, αSMA protein expression increased significantly (**P < 0.001) in HCF controls; however, the p38 inhibitors decreased this expression in T1-stimulated cells. p38inhns (063 and Birb) decreased significantly (*P < 0.05) as compared with HCF-T1 samples. β-actin was used as the loading control.
wounded cornea from transitioning to myofibroblasts, then we may be able to obtain scar-free healing. In the present study, with the addition of p38inh to T1-stimulated HCF, fibrotic protein or gene (cFN, TSP1, and Col III) expression was significantly reduced, similar to what was observed with a SMA, indicating that p38inh has the potential to decrease fibrotic protein deposition in ECM during corneal wound healing, and thus preventing the conversion of stromal cells to myofibroblasts.

Fibrosis is defined as a persistent scar. In the cornea, when fibrosis occurs, the cornea becomes opaque, and, at present, the only treatment is transplantation. Interestingly, it has been reported that T1 expression increased in an animal model and patients with lung fibrosis,9,25,50–52 indicating it is possible that T1 has a central role in persistent fibrosis. In addition, myofibroblasts have been found to persist in fibrotic organs and tissues, as well as to make fibrotic proteins.17,46,53–56 Therefore, we hypothesized that blocking the p38 pathway would change the myofibroblasts back to fibroblasts, because p38inh has the ability to inhibit both exogenous and endogenous T1 in HCF (see Figs. 3–5). Our data indicate that with the addition of T1 to HCF in culture, a SMA protein expression increased, thus transitioning the HCF to myofibroblasts and once the T1 was removed, p38inh enhanced the decrease in a SMA protein levels. When these myofibroblasts remained in T1 medium, the a SMA protein levels continued to increase with time, thus mimicking the fibrotic in vivo state (Fig. 9B). Excitingly, with the addition of p38inh to these cells, a SMA protein levels decreased similarly to those cells in p38inh-only medium (Fig. 8). These data were confirmed by morphology and localization studies (Fig. 9A, 9C), indicating that blocking the p38 pathway can change human corneal myofibroblast phenotypes in culture, which would be potentially important for the treatment of opaque corneas.

In conclusion, the results from this study indicate that HCF transition to myofibroblasts is mainly through the p38 pathway.

Figure 7. The role of the p38 inhibitor on cFN and Col III expression. HCF were cultured in the following medium: (1) BM, EMEM only, (2) T1, BM + T1 (2 ng/mL), (3) p38inh, BM + p38inh (10 μM), and (4) T1 + p38inh, BM + T1 + p38inh, and harvested for (A) WB analysis for cFN protein expression and (B) qRT-PCR analysis of Col III mRNA expression. (A) cFN protein expression increased significantly in T1-stimulated HCF (A, lane 2; *P < 0.05) as compared with BM control. The addition of p38inh to T1 samples significantly inhibited T1’s role on the cFN protein levels (A, lane 4; *P < 0.05). A representative blot from one of three independent experiments is shown (inset). β-actin was used as the loading control. (B) Col III mRNA levels were upregulated by T1 (*P < 0.05); however, the presence of p38inh efficiently inhibited T1’s role in Col III mRNA expression (*P < 0.05).

Figure 8. Analysis of a SMA expression in HCF cultured as follows: (1) HCF, RM (EMEM + 10% fetal bovine serum), (2) T1, RM + 2 ng/mL T1 (for 3 days), or (3) RM + T1 for 5 days, then RM + p38inh for D1, D2, or D3. (A) Graph of a SMA WB corrected per lane with internal control (β-actin). a SMA expression in T1 samples was significantly different than in untreated cells (HCF; P < 0.0001). On D2, a SMA expression in p38inh-treated cells decreased significantly as compared with cells in RM (*P < 0.05). (Inset) Representative WB image from one of three independent experiments. (B) Graph of relative a SMA mRNA corrected with β-actin, an internal control. a SMA mRNA data agree with protein data (A).
and blocking the p38 pathway can reduce and reverse αSMA levels. In addition, other fibrotic-associated proteins were inhibited in T1-stimulated HCF. Therefore, the p38 inhibitor is a potential therapeutic tool for human corneal fibrosis prevention/treatment, because it controls myofibroblast formation in human corneal cells while leaving other functions of T1 unaffected.

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