WNT10B Enhances Proliferation through β-Catenin and RAC1 GTPase in Human Corneal Endothelial Cells*

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Background: Adult human corneal endothelial cells are G1-arrested, often necessitating transplantation in patients with endothelial dysfunction.
Results: WNT10B enhances proliferation in human corneal endothelial cells.
Conclusion: Modulation of the WNT10B pathway could be used to treat vision loss because of corneal endothelial dysfunction.
Significance: Modulation of the WNT10B pathway may provide a means of addressing the impending increase in donor cornea demand.

The cornea is the anterior, transparent tissue of the human eye that serves as its main refractive element. Corneal endothelial cells are arranged as a monolayer on the posterior surface of the cornea and function as a pump to counteract the leakiness of its basement membrane. Maintaining the cornea in a slightly dehydrated state is critical for the maintenance of corneal transparency. Adult human corneal endothelial cells are G1-arrested, even in response to injury, leading to an age-dependent decline in endothelial cell density. Corneal edema and subsequent vision loss ensues when endothelial cell density decreases below a critical threshold. Vision loss secondary to corneal endothelial dysfunction is a common indication for transplantation in developed nations. An impending increase in demand for and a current global shortage of donor corneas will necessitate the development of treatments for vision loss because of endothelial dysfunction that do not rely on donor corneas. Wnt ligands regulate many critical cellular functions, such as proliferation, making them attractive candidates for modulation in corneal endothelial dysfunction. We show that WNT10B causes nuclear transport and binding of RAC1 and β-catenin in human corneal endothelial cells, leading to the activation of Cyclin D1 expression and proliferation. Our findings indicate that WNT10B promotes proliferation in human corneal endothelial cells by simultaneously utilizing both β-catenin-dependent and -independent pathways and suggest that its modulation could be used to treat vision loss secondary to corneal endothelial dysfunction.

The cornea consists of the epithelium, stroma, and endothelium and is the anterior, transparent tissue of the human eye that serves as its main refractive element. The corneal endothelium plays a critical role in maintaining corneal transparency, which is necessary for sharp vision by regulating corneal hydration through its pump function (1, 2). Adult human corneal endothelial cells (CECs)2 are arranged as a monolayer on the posterior surface and are G1-arrested (3, 4). As a result, there is an age-dependent decrease in CEC density that can be accelerated further by infection or injury. Corneal edema with subsequent vision loss results when the density decreases below a critical threshold of ~500 cells/μm2. The current standard of care for severe vision loss secondary to endothelial dysfunction is corneal transplantation in the form of endothelial keratoplasty. Fortunately, there currently is no shortage of donor corneas in developed nations, but the demand is expected to increase because of the increasing age of the general population and an increasing number of patients undergoing intraocular surgery. Moreover, there are ~8 million patients worldwide who have not been treated because of a local lack of donor corneas (5). Investigation of treatment strategies that do not rely on donor corneas, such as stimulating endogenous regeneration, may hold a key to addressing the current global shortage of donor corneas, which will likely be exacerbated further by the impending increase in donor cornea demand in developed nations.

A number of soluble factors have been reported to induce cell proliferation in the corneal endothelium, including IL-1β, a key mediator of inflammation in the cornea (6–8). In our previous studies, we reported that IL-1β stimulation led to FGF2 expression, resulting in enhanced proliferation in human and rabbit CECs (9–14). However, IL-1β and FGF2 have also been shown to be key mediators of endothelial-mesenchymal transition (EnMT) that can lead to retrocorneal membrane formation when lasting restoration of vision is no longer possible because of fibrosis, thereby making their use potentially dangerous (6, 10, 15). For a candidate to be clinically useful for treatment of vision loss secondary to endothelial dysfunction, it has to be able to promote proliferation in human CECs without inducing fibrosis.

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2 The abbreviations used are: CEC, corneal endothelial cell; EnMT, endothelial-mesenchymal transition; IL-1ra, interleukin-1 receptor antagonist; SFM, serum-free medium; GTPγS, guanosine 5’-3-O-(thio)triphosphate; PCNA, proliferating cell nuclear antigen; ANOVA, analysis of variance; XAV, XAV939; DI, domain inhibitor; RA, RHOC activator; Y, Y27632; NSC, NSC23766; sFRP, secreted Frizzled-related peptide; NT, non-targeting; Veh C, vehicle control; Fasc, fasapacycin; Pos C, positive control.

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26752 JOURNAL OF BIOLOGICAL CHEMISTRY
The Wnt family of ligands regulates many processes that are critical for development and cellular homeostasis (16–20). Wnt signaling can be broadly classified into canonical (β-catenin-dependent) and non-canonical (β-catenin-independent) pathways. In the canonical pathway, binding of Wnt ligand to its receptor complex leads to nuclear transport of β-catenin, during which it interacts with the LEF-1/TCF family of transcription factors to activate transcription of target genes, including those critical for cell proliferation, such as Cyclin D1 and c-myc (16, 21–27). During eye development, the canonical Wnt signaling plays an important role in differentiation of the lens epithelium and lens fiber cells (28, 29). The non-canonical pathway does not signal through β-catenin and includes the planar cell polarity and Ca\(^{2+}\) pathways (30, 31). These pathways have been shown to regulate development, including extension movements during vertebrate gastrulation (32) and cell migration (33, 34). We have reported previously that WNT5A signals through the non-canonical pathway in human CECs and promotes migration through activation of CDC42 and inhibition of RHOA (35).

The RhoGTPase family members RAC1, RHOA, and CDC42 modulate various cellular functions through cycling between active GTP-bound and inactive GDP-bound forms (36, 37). It has been reported that these proteins regulate planar cell polarity in Drosophila (38) and convergent extension movement in Xenopus (32, 39). However, RAC1 has also been reported to be involved in the canonical pathway through modulation of β-catenin nuclear transport (40) and activation of β-catenin/TCF-mediated transcription of Cyclin D1 (41). The role of RAC1 and β-catenin signaling in proliferation of human CECs has not been reported previously.

In this study, we investigated the downstream targets of IL-1β capable of driving proliferation in human CECs. Our results show that IL-1β activates WNT10B expression and promotes proliferation in human CECs. WNT10B signals through Disheveled and induces independent nuclear transport of β-catenin during which they form a complex to activate expression of Cyclin D1, leading to proliferation of human CECs. We provide evidence that WNT10B signals simultaneously through β-catenin-dependent and -independent pathways to enhance proliferation in human CECs, and our results suggest that modulation of WNT10B could be used to treat vision loss secondary to corneal endothelial dysfunction.

Experimental Procedures

Reagents—FGF2 was purchased from Cell Signaling Technology (Danvers, MA). Secreted Frizzled-related peptide (sFRP) 1 (42) was purchased from Fitzgerald Industries International (Acton, MA). Anti-JUN antibody, Disheveled-PDZ domain inhibitor II (25 µM), sulfasalazine (an inhibitor of IκB degradation, 2 mM), Disheveled-PDZ domain inhibitor II (25 µM), XAV939 (20 µM), RHOA activator (1 µg/ml), Y27632 (a Rho-associated coiled kinase inhibitor, 10 µM), ML141 (a CDC42 inhibitor, 10 µM), or NSC23766 (a RAC1 inhibitor, 100 µM). All experiments included a vehicle control (0.1 mM EDTA and 0.5% CHAPS in PBS or dimethyl sulfoxide) sample.

ChIP Assay—FGF2-starved human CECs were first pretreated with sulfasalazine or SB203580 for 2 h and then stimulated with IL-1β or vehicle control for 10 min. The cells were then maintained in SFM for 6 h prior to isolating protein-DNA complexes using anti-NF-κB (p65) and AP-1 (c-JUN) antibodies. Binding of NF-κB and AP-1 to the WNT10B promoter was evaluated by PCR using the following primer pairs: NF-κB site 1, 5’-CTGTCTAAGGTATCCCATGG-3’ (forward) and 5’-TGAATCTACCTTGTTGACGCC-3’ (reverse); NF-κB site 2, 5’-CTACTCGCCCTTGTGATGACACC-3’ (forward) and 5’-CTCTGTCTTCAAACCTGCCC-3’ (reverse); and NF-κB site 3, 5’-ACTCCCTGAGGTGTAAAGACC-3’ (forward) and 5’-CTCCAGGGGTTAAGCTTT-3’ (reverse). Standard PCR conditions were as follows: 5 min at 94 °C, followed by 33 cycles of 30 s at 94 °C, 30 s at 55 °C, 30 s at 72 °C, and a final extension for 2 min at 72 °C. Annealing temperature was adjusted depend on the PCR primer. PCR products were separated by 1.5% agarose gel electrophoresis and visualized by ethidium bromide staining.

Isolation and Culture of Primary Human CECs—Isolation and culture of primary human CECs were performed according to previously published protocols (12, 47) with minor technical modifications. Briefly, corneas were removed from Optisol and washed several times with OptiMEM I (Gibco-BRL) containing...
50 μg/ml gentamicin. The Descemet membrane and endothelium complex was stripped from corneas and treated with 0.2% collagenase type II and 0.05% hyaluronidase (Worthington Biochemical, Lakewood, NJ) for 3 h at 37 °C. After centrifugation, the primary cells were resuspended in culture medium (OptiMEM I supplemented with 5% fetal bovine serum, 10 μM Y27632, 0.5 μg/ml R-spondin, and 10 μM SB431542 (TGF-β receptor inhibitor kinase inhibitor VI) (48) and plated on 24-well tissue culture plates precoated with FNC Coating Mix (Biological Research Faculty and Facility, Inc., Ijamsville, MD) and laminin (Sigma). For subculture, confluent primary cultures were treated with 0.05% trypsin and 5 mM EDTA in PBS for 10 min. Second- or third-passage human CECs were used for all experiments. For serum starvation, the culture medium was changed to OptiMEM I, and cultures were maintained for 24 h. In some experiments, pharmacologic inhibitors were used in the presence of IL-1β (5 ng/ml), FGF2 (10 ng/ml), WNT10B (200 ng/ml), WNT5A (300 ng/ml), or WNT3A (50 ng/ml) stimulation: IL-1ra (50 ng/ml), AZD4547 (an FGF receptor inhibitor, 0.1 μM), sFRP (3 μg/ml), Disheveled-PDZ domain inhibitor II (25 μM), XAV939 (20 μM), RHOA activator (1 μg/ml), Y27632 (10 μM), ML141 (10 μM), or NSC23766 (100 μM). Vehicle control (0.1 mM EDTA and 0.5% CHAPS in PBS or dimethyl sulfoxide) was included for all experiments.

In Vitro Proliferation Assays—Primary human CECs were used for all cell proliferation assays. The 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide assay was used to assess proliferation as described previously (10). Briefly, cells were seeded in 96-well tissue culture plates at a concentration 4 × 10^5 cells/well. When cells reached ~70% confluence, the medium was changed to OptiMEM I for serum starvation and maintained for 24 h. The serum-starved cells were then maintained for 24 h under each culture condition. Afterward, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (50 μg/ml) was added, and the culture was maintained for another 2 h at 37 °C. The 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (50 μg/ml) was added, and the culture was maintained for another 2 h at 37 °C. The 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide assay was used to assess proliferation as described previously (10).

RHOA activation was determined by Western blot analysis using total cell lysates. Results were normalized to the absorbance of lysate from untreated SFM-maintained cells and expressed as a percentage of this value for the comparison of RHOA activity (level of GTP-bound RHOA). Nuclear lysate protein concentrations were assessed with the Bradford protein assay system (Bio-Rad), and lysates were then used in coimmunoprecipitation experiments.

RHOA GTPase Activation Assay—RHOA activation was quantified as described previously (35) by measuring the amounts of RHOA-GTP using the ELISA-based G-LISA™ RHOA activation biochem assay kit (Cytoskeleton Inc.). Active RHOA-GTP from lysates bound to the GTP-Rho binding domain of the Rho effector protein was detected by primary anti-RHOA antibody and HRP-conjugated secondary antibody. Results were normalized to the absorbance of lysate from SFM-maintained cells and expressed as a percentage of this value for the comparison of RHOA activity (level of GTP-bound RHOA) in different cell lysates.

Ex Vivo Proliferation Assay—Whole human corneas not suitable for transplantation were purchased from Vision Share (Ann Arbor, MI) and cut into three equal sections. The endothelium of each piece was injured with a small pipette tip measuring ~0.5 mm in diameter, resulting in a linear wound ~0.5 mm wide. The corneal pieces were then placed endothelial side up in individual wells of a 24-well tissue culture plate. The pieces were maintained with OptiMEM containing WNT10B with or without inhibitors for 72 h at 37 °C in a 5% carbon dioxide, humidified atmosphere. Pharmacologic inhibitors were used in the presence of WNT10B (200 ng/ml) stimulation: sFRP (3 μg/ml), Disheveled-PDZ domain inhibitor II (25 μM), XAV939 (20 μM), NSC23766 (100 μM), or fasaplysin (0.5 μM). For siRNA knockdown in ex vivo human corneal endothelium, the Accell SMARTpool system was used. 36 h after transfection
WNT10B Enhances Proliferation in Corneal Endothelial Cells

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Corneal endothelial cells (CECs) are a monolayer of cells located at the posterior corneal surface. They are crucial for maintaining the long-term integrity of the corneal endothelium. WNT10B, a member of the Wnt family, has been shown to play a role in regulating proliferation and differentiation of various cell types. In this study, the authors investigated the role of WNT10B in proliferation of human corneal endothelial cells.

**Figure 1.** IL-1β stimulation enhances proliferation and activates expression of WNT10B in human CECs. A, treatment of primary human CECs with IL-1β (+) increased the relative proliferation rate compared with Veh C (152 ± 3.8 versus 100 ± 7.5, p < 0.01). IL-1β-dependent proliferation was abolished by cotreatment with IL-1ra (112 ± 5.4 versus 152 ± 3.8, p < 0.01). Cotreatment with AZD4547 (AZD), a FGF receptor 1–3 antagonist, attenuated IL-1β-enhanced proliferation (130 ± 4.3 versus 152 ± 3.8, **p < 0.01), and it abolished FGF2-enhanced proliferation in human CECs (104 ± 2.4 versus 173 ± 7.1; ***, p < 0.001). Treatment with AZD4547 alone did not change the proliferation rate compared with vehicle control (98 ± 3.7 versus 100 ± 7.5, p > 0.05). The relative proliferation rate was greater in IL-1β-stimulated CECs that were cotreated with AZD4547 compared with those cotreated with IL-1ra (130 ± 4.3 versus 112 ± 5.4; p < 0.01). One-way ANOVA, F(6, 35) = 186, p < 0.00001, n = 6/sample. Tukey’s post-hoc test, HSD(0.05) = 9.4 and HSD(0.01) = 11.3, where HSD is honestly significant difference. B, treatment of human CECs with IL-1β (+) activated expression of WNT10B, and IL-1β-dependent WNT10B expression could be abolished with IL-1ra (ra) cotreatment. IL-1β-dependent expression of WNT10B could be attenuated by cotreatment with either the NF-κB inhibitor sulfasalazine (Sul) or the AP-1 inhibitor SB203580 (SB). Cotreatment with both inhibitors abolished IL-1β-induced WNT10B expression. There was no effect on p65 and JUN protein levels by cotreatment with the inhibitors. Transfection with p65 siRNA and JUN siRNA resulted in corresponding knockdown of p65 and JUN protein levels in human CECs. Transfection with either p65 or JUN siRNA resulted in attenuation of IL-1β-dependent WNT10B expression, whereas transfection with both p65 and JUN siRNA severely decreased WNT10B expression. Transfection with NT control siRNA had no effect on p65, JUN, and WNT10B expression in human CECs. β-actin was used as a loading control. C, the expected PCR product sizes for NF-κB and AP-1 binding site were as follows: NF-κB site 2 (N1), 260 bp; NF-κB site 2 (N2), 147 bp; NF-κB site 3 (N3), 140 bp; AP-1 site 1 (A1), 380 bp; and AP-1 site 2 (A2), 204 bp. IL-1β, but not vehicle control treatment, resulted in binding of NF-κB and AP-1 to the WNT10B promoter in human CECs. NF-κB binding could be antagonized by pretreatment with sulfasalazine but not SB203580, and AP-1 binding could be antagonized with pretreatment with SB203580 but not sulfasalazine. Anti-histone H3 antibody was used as a positive control (+), and normal rabbit IgG was used as a negative control (−).

Statistical Analysis—One-way analysis of variance (ANOVA) was performed to compare means within groups, and post-hoc Tukey’s honest significant difference tests were done to perform pairwise comparisons between means within groups.

With 1.5 μM Accell SMARTpool of siRNA targeting Disheveled 2 or Cyclin D1 (Dharmacon) in Accell Delivery media, the medium was changed to Opti-MEM with WNT10B and maintained for another 72 h. Accell non-targeting control siRNA (Dharmacon) was used as a negative control. After 72 h, the corneal pieces were fixed for 20 min in 1 ml of 4% paraformaldehyde and rinsed three times with PBS. They were then permeabilized using 0.5% Triton X-100 (Sigma-Aldrich) in PBS for 30 min at room temperature, followed by blocking with 4% bovine serum albumin (Sigma-Aldrich) in PBS for 1 h at room temperature. The tissues were incubated for 2 h at room temperature with both mouse anti-human proliferating cell nuclear antigen (PCNA) IgG (Sigma-Aldrich) or mouse anti-human phosphohistone H3 IgG (EMD Millipore) and rabbit anti-human ZO-1 IgG (Invitrogen) diluted 1:100 in blocking buffer. Each piece was rinsed in PBS three times for 10 min, followed by incubation with FITC-conjugated goat anti-mouse IgG (Sigma-Aldrich) and Rhodamine-conjugated donkey anti-rabbit IgG (Genetex) diluted 1:200 in blocking buffer for 2 h at room temperature. Incubation of primary or secondary antibody alone was used as a negative control. The corneal pieces were then washed three times for 10 min with PBS at room temperature. Excess sclera was removed to facilitate mounting (endothelial side down) with mounting solution with DAPI on a glass-bottom microwell dish (MatTek Co., Ashland, MA). A glass coverslip was placed on the epithelial side of the corneal sections, and a 20-g weight was placed on the coverslip for 15 min to flatten the sections for imaging. Images were captured with a BZ-X700 fluorescence microscope (Keyence, Itasca, IL) using a ×40 objective lens with an aperture of 0.95. All images were captured at room temperature on sections mounted with Vectashield with DAPI mounting medium (Vector Laboratories, Burlingame, CA) and analyzed with BZ-X analyzer software (Keyence). PCNA-positive nuclei were counted in an 85-μm² field within the wound area. Five fields were counted per section, and three different sections were used for each culture condition. The proliferation index was calculated as the number of PCNA-positive nuclei divided by the number of all nuclei within each field, and the relative proliferation rate was calculated relative to the proliferation index of the vehicle-treated cornea.
IL-1β Stimulation Leads to Cell Proliferation and WNT10B Expression in Human CECs—We have reported previously that IL-1β stimulation led to FGF-2 expression in human (9) and rabbit (10, 11) CECs, resulting in enhanced proliferation (12–14). The FGF receptor 1–3 antagonist AZD4547 attenuated, but did not completely abolish, IL-1β-induced proliferation (Fig. 1A), suggesting that IL-1β-induced proliferation was mediated by a signaling pathway distinct from FGF2 in human CECs. Stimulation of human CECs with IL-1β resulted in activation of WNT10B expression, which, in turn, could be inhibited by cotreatment with the IL-1 receptor antagonist (Fig. 1B). IL-1β-dependent WNT10B expression in human CECs could be antagonized by pretreatment with the NF-κB inhibitor sulfasalazine or the AP-1 antagonist SB203580 (Fig. 1B). Pretreatment with both sulfasalazine and SB203580 resulted in complete inhibition of IL-1β-dependent WNT10B expression (Fig. 1B). Furthermore, IL-1β stimulation of human CECs transfected with either p65 siRNA targeting NF-κB or JUN siRNA targeting AP-1 resulted in attenuation of WNT10B expression, whereas transfection with both p65 and JUN siRNA completely inhibited IL-1β-induced WNT10B expression (Fig. 1B). Corresponding knockdowns of p65 and JUN proteins were observed in p65 and JUN siRNA-transfected human CECs, and neither use of small molecule inhibitors nor transfection with a nontargeting control siRNA affected p65 and JUN protein levels. The human WNT10B promoter contains several conserved and putative NF-κB and AP-1 binding sites (49). Binding of NF-κB and AP-1 to the WNT10B promoter in IL-1β-treated human CECs was shown by chromatin immunoprecipitation assays (Fig. 1C). NF-κB binding could be antagonized by pretreatment with sulfasalazine but not SB208530, and AP-1 binding could be antagonized by pretreatment with SB208530 but not sulfasalazine (Fig. 1C). Vehicle control treatment did not result in NF-κB and AP-1 binding. Treatment with IL-1β led to expression of WNT10B in human ex vivo corneal endothelium, and combining injury with IL-1β treatment resulted in greater WNT10B expression (Fig. 1D). Neither vehicle control nor injury alone resulted in expression of WNT10B in human ex vivo corneal endothelium.

WNT10B Signaling Proceeds through the Canonical Pathway in Human CECs—β-catenin was found in the cytoplasmic fraction but not in the nuclear fraction of human CECs treated with vehicle control (Fig. 2A). Treatment of human CECs with the positive control (WNT3A) resulted in nuclear transport of β-catenin whereas treatment with the negative control (WNT5A) did not. WNT3A was shown to signal through the canonical Wnt pathway, whereas WNT5A was shown to signal through the non-canonical Wnt pathway in human CECs.3 Treatment with WNT10B resulted in nuclear transport of β-catenin (Fig. 2A). WNT10B-dependent nuclear transport of β-catenin could be inhibited by cotreatment with sFRP, the β-catenin antagonist XAV, and Disheveled-PDZ domain inhibitor (DI) (Fig. 2B). Cotreatment with RHOA activator (RA), the Rho kinase inhibitor Y27632 (Y), and the RAC1 inhibitor NSC23766 (NSC) had no effect on WNT10B-dependent nuclear transport of β-catenin (Fig. 2B). Lamin B was used as a nuclear fraction marker, and α-tubulin was used as a cytoplasmic fraction marker.

WNT10B Activates RAC1 and Inhibits RHOA in Human CECs—RhoGTPases have been reported previously to regulate cell proliferation through modulation of β-catenin signaling (40, 41, 50, 51). GTP-bound CDC42 was detected in positive control (WNT5A)-treated cells but not in vehicle control-treated human CECs (Fig. 3A). WNT10B treatment of human CECs did not result in CDC42 activation (Fig. 3A). WNT10B stimulation led to activation of RAC1 in human CECs, as shown by the presence of GTP-RAC1

3 J. G. Lee and M. Heur, unpublished data.

26756 JOURNAL OF BIOLOGICAL CHEMISTRY

VOLUME 290 • NUMBER 44 • OCTOBER 30, 2015

ASBMB
WNT10B-dependent RAC1 activation could be inhibited with cotreatment with sFRP, the RAC1 inhibitor NSC, and Disheveled-PDZ DI. Cotreatment with the β-catenin antagonist XAV, RA, and the Rho kinase inhibitor Y had no effect on WNT10B-dependent RAC1 activation in human CECs. The bar graph depicts the relative -fold difference in GTP-RAC1 levels. GTP-RAC1 was detected in Pos C-treated (WNT5A-treated) but not in Veh C-treated cells. WNT10B treatment (+) inhibited RHOA activity relative to Veh C (100 ± 3.9 versus 65 ± 2.3, p < 0.01). WNT10B-dependent inhibition of RHOA activity was abolished by cotreatment with sFRP (93 ± 4.4 versus 65 ± 2.3; *, p < 0.01) and DI (89 ± 2.5 versus 65 ± 2.3; **, p < 0.01), but cotreatment with XAV (77 ± 4.2 versus 65 ± 2.3, p > 0.05) or NSC (72 ± 5.4 versus 65 ± 2.3, p > 0.05) had no effect. Cotreatment with RA in the absence of WNT10B stimulation resulted in higher RHOA activity (154 ± 6.1 versus 100 ± 3.9, p < 0.01), and treatment with the Pos C, WNT5A, resulted in lower RHOA activity (48 ± 2.0 versus 100 ± 3.9, p < 0.01) than Veh C. One-way ANOVA, F(7, 24) = 102, p < 0.00001, n = 4/sample. Tukey’s post-hoc test, HSD[0.05] = 14.4 and HSD[0.01] = 17.3.

D, treatment with WNT10B induced Disheveled 2 and RAC1 binding. A 3.5-fold increase in anti-Disheveled 2 antibody-precipitated lysates from WNT10B-treated (+) compared with vehicle control (C)-treated human CECs. Cotreatment with sFRP, DI, and NSC inhibited WNT10B-dependent Disheveled 2 and RAC1 binding, whereas cotreatment with XAV, Y, and RA had no effect on WNT10B-dependent Disheveled 2 and RAC1 binding. The bar graph depicts the relative -fold difference in RAC1 levels. Transfection with Disheveled 2 siRNA also blocked WNT10B (W10B)-dependent binding between Disheveled 2 and RAC1, whereas NT control siRNA had no effect. IP, immunoprecipitation; IB, immunoblot.
WNT10B Enhances Proliferation in Corneal Endothelial Cells

WNT10B stimulates proliferation in human CECs—WNT10B treatment resulted in a significant increase in relative proliferation rates compared with vehicle control in human CECs (Fig. 5A). WNT10B-dependent proliferation could be inhibited by cotreatment with sFRP, Disheveled-PDZ DI, and the β-catenin antagonist XAV (Fig. 5A). Cotreatment with the negative control (WNT5A) had no effect, whereas cotreatment with the positive control (WNT3A) enhanced proliferation compared with vehicle control. Inhibition of RAC1 by cotreatment with NSC resulted in inhibition of WNT10B-dependent proliferation, whereas cotreatment with the Rho kinase inhibitor Y, RA, and the CDC42 antagonist ML141 did not have an effect (Fig. 5B). Transfection of primary human CECs with RAC1 and β-catenin siRNAs resulted in knockdown of RAC1 and β-catenin protein levels, whereas transfection with NT control siRNA did not alter RAC1 and β-catenin protein levels (Fig. 5C). RAC1 and β-catenin siRNA transfections significantly inhibited WNT10B-dependent proliferation in primary human CECs (Fig. 5D).

Cyclin D1 Mediates WNT10B-dependent Cell Proliferation in Human CECs—Cyclin D1 has been reported previously as a transcriptional target of Wnt/β-catenin signaling (25, 55), and RAC1 has also been reported previously to activate Cyclin D1 expression through β-catenin in other cell types (41, 52). WNT10B stimulation increased Cyclin D1 expression compared with vehicle control in human CECs (Fig. 6A). WNT10B-dependent activation of Cyclin D1 expression could be inhibited by cotreatment with sFRP and Disheveled-PDZ DI and, to a lesser extent, by cotreatment with the β-catenin antagonist XAV and the RAC1 inhibitor NSC (Fig. 6A). Cotreatment with the Rho kinase inhibitor Y and RA did not have an effect on WNT10B-dependent activation of Cyclin D1 expression. Transfection of human CECs with Cyclin D1 siRNA resulted in a corresponding knockdown of Cyclin D1 protein levels in WNT10B-treated human CECs, whereas transfection with non-targeting control siRNA had no effect (Fig. 6B). Cyclin D1 (CCND1) siRNA transfection abolished WNT10B-dependent proliferation in human CECs, but transfection with non-targeting control siRNA had no effect compared with vehicle control (Fig. 6C). Transfection with NT control siRNA had no effect on relative proliferation rates.

WNT10B Promotes Proliferation in Human CECs ex Vivo—To further investigate the role of WNT10B signaling in human CEC proliferation, the proliferative response of endothelial cells to wounding in human ex vivo corneal endothelium was examined. PCNA and zonula occludens 1 (ZO-1) were used as

lysates precipitated with anti-Disheveled 2 antibody (Fig. 3D). This is consistent with previous reports describing binding between Disheveled with RAC1 in other cell types (52–54). WNT10B-dependent binding of Disheveled 2 and RAC1 could be inhibited by cotreatment with sFRP, Disheveled-PDZ DI, or NSC. Cotreatment with XAV, RA, or Y had no effect on WNT10B-dependent binding between Disheveled and RAC1 (Fig. 3D). Transfection of human CECs with Disheveled 2 siRNA resulted in the corresponding knockdown of Disheveled 2 protein and inhibited WNT10B-dependent binding between Disheveled 2 and RAC1 (Fig. 3D). Transfection with non-targeting control siRNA had no effect on Disheveled 2 protein levels or WNT10B-dependent binding between Disheveled and RAC1 (Fig. 3D).

WNT10B Activates Nuclear Transport and Binding of RAC1 and β-Catenin—It has been reported previously that RAC1 plays a role in the nuclear transport of β-catenin (40, 41, 52). RAC1 and β-catenin bound to RAC1 were present in the nuclear fraction of WNT10B but not vehicle control-treated human CECs (Fig. 4A). WNT10B-dependent nuclear transport of RAC1 and its association with β-catenin were blocked by cotreatment with sFRP, Disheveled-PDZ DI, and the RAC1 inhibitor NSC, whereas cotreatment with the Rho kinase inhibitor Y and RA did not have an effect (Fig. 4A). Cotreatment with the β-catenin antagonist XAV did not interfere with nuclear transport of RAC1, but it was not complexed with β-catenin in the nucleus (Fig. 4A). Although RAC1 was present in the cytoplasmic fraction under all conditions, it was not bound to β-catenin in the cytoplasm (Fig. 4B). Lamin B was used as a nuclear fraction marker, and α-tubulin was used as a cytoplasmic fraction marker.

FIGURE 4. WNT10B induces nuclear transport and binding of RAC1 and β-catenin. A, an approximate 8-fold increase in RAC1 level along with an approximate 6-fold increase β-catenin level were detected in the nuclear fractions precipitated with RAC1 antibody in WNT10B-treated (+) but not in Veh C-treated human CECs. Cotreatment with sFRP, Disheveled-PDZ DI, and the RAC1 inhibitor NSC blocked WNT10B-dependent nuclear transport of RAC1 and its interaction with β-catenin. Inhibition of β-catenin with XAV treatment did not interfere with nuclear transport of RAC1, but β-catenin was not found in the nucleus complexed to RAC1. Cotreatment with RA or the Rho kinase inhibitor Y had no effect on WNT10B-dependent RAC1 nuclear transport and its interaction with β-catenin. The bar graphs depict the relative -fold difference in RAC1 and β-catenin levels. Lamin B and α-tubulin were used as nuclear and cytoplasmic markers, respectively. IP, immunoprecipitation; IB, immunoblot. B, RAC1 but not β-catenin complexed to RAC1 was detected in the cytoplasm under all treatment conditions.
markers for proliferation and CECs, respectively, in human ex vivo corneas. Exogenous WNT10B could not promote proliferation in CECs of human corneas ex vivo when the endothelium was not wounded (data not shown). WNT10B, but not vehicle control (Veh C), induced expression of PCNA in CECs in wounded endothelium of ex vivo human corneas (Fig. 7A). WNT10B-dependent expression of PCNA in human CECs ex vivo could be antagonized by cotreatment with sFRP, Disheveled-PDZ DI, the β-catenin antagonist XAV, NSC, and the CDK4-Cyclin D1 complex inhibitor fascaplysin (Fasc) (Fig. 7A). Quantification of PCNA expression in ex vivo human corneal endothelium showed significantly increased relative proliferation rates in response to WNT10B treatment (Fig. 7B). This, in turn, could be inhibited significantly by cotreatment with sFRP (p < 0.01), Disheveled-PDZ DI (p < 0.01), XAV (p < 0.01), NSC (p < 0.01), and Fasc (p < 0.01), and there was no statistically significant difference between the control group and the WNT10B group (p = 0.05). One-way ANOVA, F(4, 40) = 34.2, p < 0.00001, n = 9/sample. Tukey’s post-hoc test, HSD(0.05) = 14.3 and HSD(0.01) = 17.5.
WNT10B Enhances Proliferation in Corneal Endothelial Cells

significant difference in PCNA expression between Veh C treatment and WNT10B + antagonist (sFRP, DI, XAV, NSC, and Fasc) treatment (p > 0.05, Fig. 7B). CCND1 siRNA transfection of ex vivo human corneal endothelium resulted in a knockdown of WNT10B-induced Cyclin D1 protein levels (Fig. 7C). Transfection with Disheveled 2 (DVL2) siRNA led to knockdown of DVL2 protein levels and inhibition of WNT10B-dependent Cyclin D1 expression (Fig. 7C). NT control siRNA transfection did not alter Cyclin D1 and Disheveled 2 protein levels. Transfection with Cyclin D1 and Disheveled 2 siRNA abolished PCNA expression in wounded ex vivo corneal endothelium treated with WNT10B, whereas non-targeting siRNA transfection did not affect PCNA expression (Fig. 7D). Cyclin D1 and Disheveled 2 siRNA transfection also significantly reduced relative proliferation rates in wounded ex vivo corneal endothelium treated with WNT10B (Fig. 7E). Transfection with non-targeting siRNA had no effect. To further evaluate cell cycle reentry of human ex vivo CECs, phosphohistone H3 was used as another marker. Treatment with WNT10B following injury led to phosphorylation of histone H3 whereas treatment with vehicle control did not (Fig. 7F).

Discussion

Unlike other cells, in which proliferation plays a critical role in wound healing, endothelial cells in the adult human cornea are G1-arrested and do not proliferate even in response to injury (3, 4). This unusual feature of adult human CECs, in the setting of injury or infection, can lead to severe vision loss secondary to corneal edema. The current standard of care for severe vision loss secondary to endothelial dysfunction is corneal transplantation in the form of endothelial keratoplasty. There is currently a global shortage of donor corneas that will likely be exacerbated by an impending increase for donor corneas in developed nations. One method of addressing this issue is to develop strategies for treatment of vision loss because of endothelial dysfunction that do not rely on donor corneas. Tissue engineering approaches using human CECs derived from in vitro expansion methods are being investigated (56–61). However, these approaches have been hampered by fibroblastic transformation and uncertainty regarding the endothelial phenotype in cultured cells. Approaches targeting specific signaling pathways in human CECs or relying on alternative starting material hold the promise of generating a potentially unlimited supply of transplantable endothelium (62–65). All new approaches will have to ensure the endothelial phenotype and verify that the derived endothelial cells have not undergone mesenchymal transition to ensure safety and efficacy.

We have reported previously that IL-1β-induced FGF2 enhances proliferation and migration in rabbit and human CECs (9, 11–14). However, FGF2 has also been reported as a key mediator for ENMT, leading to retrocorneal fibrous membrane formation (13, 15, 66–68). This led us to investigate other downstream targets of IL-1β signaling in human CECs that could enhance cell proliferation without inducing fibrosis. Attenuation, but not complete inhibition, of cell proliferation by AZD4547 (Fig. 1A) strongly suggested the presence of another signaling pathway distinct from FGF2 that could drive proliferation in human CECs. Previous reports of a WNT10B promoter containing several conserved AP-1 and NF-κB sites (49) in conjunction with AP-1 and NF-κB being activated by IL-1β (9, 11, 35) led us to investigate the role of WNT10B in IL-1β-dependent proliferation in human CECs. As anticipated, IL-1β-dependent WNT10B expression proceeds through AP-1 and NF-κB activation in human CECs (Fig. 1, B and C). Also as anticipated, injury could not induce expression of WNT10B in human ex vivo CECs, but it could augment it in the presence of IL-1β (Fig. 1D). Nuclear transport of β-catenin indicated that WNT10B signals through the canonical pathway in human CECs (Fig. 2), similar to what has been reported for other cell types (16, 21, 69, 70). WNT3A was another potential candidate because it can induce nuclear transport of β-catenin in human CECs in culture, but it was not expressed in human CECs in response to IL-1β stimulation.3
Previous studies have reported that Disheveled 2 is required for the activation of RAC1 (32, 53) and that activated RAC1 regulates nuclear localization of β-catenin and β-catenin/TCF-dependent transcription (40, 52). Our results indicate that Disheveled 2 is critical for WNT10B signaling and occupies the nexus controlling both β-catenin nuclear transport and regula-
WNT10B Enhances Proliferation in Corneal Endothelial Cells

Antagonizing Disheveled 2 activity with Disheveled-PDZ D1 resulted in inhibition of WNT10B-dependent nuclear transport of β-catenin (Fig. 2B), activation of RAC1 (Fig. 3B), and inhibition of RHOA (Fig. 3C). Inhibition of Disheveled 2 activity by either DI or siRNA knockdown led to an inhibition of WNT10B-induced binding between Disheveled 2 and RAC1 (Fig. 3D). In contrast to previous reports, activated RAC1 does not appear to be required for nuclear transport of β-catenin in human CECs, as evidenced by RAC1 inhibition with NSC23766 having no effect on WNT10B-dependent nuclear transport of β-catenin in human CECs (Fig. 2B). Moreover, β-catenin does not appear to play a role in RAC1 signaling in human CECs, as evidenced by inhibition of β-catenin with XAV939 having no effect on WNT10B-dependent activation (Fig. 3B) and nuclear transport (Fig. 4A) of RAC1. Although the nuclear transport of β-catenin and RAC1 are independent of one another, both are required for WNT10B-dependent proliferation in human CECs, as evidenced by XAV939 and NSC23766 independently being able to abolish WNT10B-dependent proliferation (Fig. 5), likely through attenuation of Cyclin D1 expression (Fig. 6A). It is interesting to note that, although WNT10B inhibited RHOA activity (Fig. 6C), neither modulation of RHOA by inhibition with Y27632 nor activation with RHOA activator had any effect on WNT10B-dependent proliferation in human CEC (Fig. 5B). This suggests that RHOA plays a very minor role in WNT10B-dependent proliferation in human CECs. Downstream of β-catenin and RAC1, WNT10B-dependent proliferation in human CECs proceeds through Cyclin D1. To address the concern over nonspecific effects of small molecule inhibitors, we confirmed the roles of RAC1 and β-catenin in WNT10B-induced CEC proliferation using siRNA-mediated knockdown of RAC1 and β-catenin (Fig. 5D).

WNT10B signals simultaneously through canonical and non-canonical Wnt pathways in human CEC proliferation. Treatment with either β-catenin antagonist XAV939 or NSC23766 substantially attenuated WNT10B-dependent expression of Cyclin D1 (Fig. 6A), and transfection of human CEC with Cyclin D1 siRNA resulted in knockdown of WNT10B-dependent Cyclin D1 expression (Fig. 6B) and inhibition of WNT10B-dependent proliferation (Fig. 6C) supporting the critical role of Cyclin D1.

To address the concern that the WNT10B-dependent proliferation could be the result of a culturing artifact, we used ex vivo human corneas to investigate the role of WNT10B signaling in CEC proliferation. The ability of WNT10B to stimulate proliferation in human CECs was comparable under ex vivo (Fig. 7) and in vitro (Figs. 5 and 6) conditions. Moreover, the results of vehicle control treatment of wounded ex vivo corneas showed a negligible number of CECs with nuclear expression of PCNA, consistent with previous reports (4, 71). Concerns over nonspecific effects of small molecule inhibitors were addressed by verifying the results from small molecule inhibitor experiments by utilizing siRNA knockdown of Cyclin D1 and Disheveled 2 in ex vivo corneal endothelium (Fig. 7, D and E). To address the concern that PCNA expression could be secondary to DNA damage rather than cell cycle reentry, we used phosphohistone H3 as an additional marker. Phosphorylation of histone H3 in wounded human ex vivo CECs treated with WNT10B indicates that they have entered mitosis (Fig. 7F). Taken together, these results suggest that modulating WNT10B signaling could play an important role in promoting endothelial cell proliferation in patients with vision loss because of endothelial dysfunction.

Although Cyclin D1 plays a crucial role in WNT10B-mediated proliferation, siRNA targeting of RAC1 and β-catenin compared with Cyclin D1 resulted in a greater inhibition of WNT10B-induced proliferation in primary human CECs (Figs. 5D and 6C). Although this strongly suggests that Cyclin D1 is not the only downstream target of WNT10B in promoting proliferation in primary human CECs, the results are slightly different in ex vivo human corneas. Targeting Disheveled 2, RAC1, and Cyclin D1 had similar effects on inhibition of proliferation in wounded ex vivo endothelial cells that were treated with WNT10B (Fig. 7). This highlights the notion that corneal endothelial cell behavior is, to some extent, dependent on their local environment.

The apparent minor role of RHOA in our results appears to contradict a previous report that showed inhibition of RHOA activity with the Rho kinase inhibitor Y27632 induced cyclin D expression and proliferation in monkey CECs (72). The differences in results could be rooted in differences in species and culture conditions causing different components, e.g. RAC1, RHOA, and β-catenin, to play different roles in mediating cell proliferation. This strongly suggests that the source and culturing conditions for corneal endothelial cells significantly influence their behavior.

With investigations of alternative approaches in very preliminary stages, a major concern that needs to be addressed is the potential induction of endothelial EnMT and its undesirable sequela. Although both IL-1β and FGF2 can induce proliferation of human CECs, they also induce EnMT, making their use potentially dangerous because of fibrosis. Future studies investigating the potential induction of EnMT by WNT10B and other targets that can enhance proliferation are needed.
Enhancing endothelial pump function through cell proliferation in vivo for patients with endothelial dysfunction not only avoids the postoperative morbidity associated with cornea transplants, but it also effectively enlarges the cornea donor pool by making more donor corneas available for other causes of corneal blindness. In summary, WNT10B activates RAC1 and β-catenin, resulting in induction of Cyclin D1 expression and subsequent proliferation in human CECs (Fig. 8).

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References
WNT10B Enhances Proliferation in Corneal Endothelial Cells


