VIP and VIP Gene Silencing Modulation of Differentiation Marker N-Cadherin and Cell Shape of Corneal Endothelium in Human Corneas Ex Vivo

Shay-Whey M. Koh,1,2 Krish Chandrasekara,3 Cara J. Abbondandolo,1 Timothy J. Coll,1 and Allan R. Rutzen1

PURPOSE. Vasoactive intestinal peptide (VIP) is expressed by corneal endothelial (CE) cells and is present in the aqueous humor, which bathes CE cells in vivo. This study demonstrated the role of CE cell VIP in maintaining the expression level of a CE differentiation marker, N-cadherin, and the hexagonal cell shape.

METHODS. To determine the most effective VIP concentration, bovine corneoscleral explants were treated with 0 (control) and 10−12 to 10−6 M VIP. Paired human corneas (nine donors) from an eye bank were used as control; the other corneas were treated with VIP. To silence endogenous VIP, paired fresh human donor corneas (from seven cadavers) were transduced with VIP shRNA or the control lentiviral particles and then bisected/quartered for quantitative analysis by semiquantitative RT-PCR (for mRNA) and Western blot analysis/immunocytochemistry (for protein), whereas alizarin red S staining revealed CE cell shape.

RESULTS. VIP concentration dependently increased bovine CE cell N-cadherin mRNA levels, with the maximal effect observed between 10−10 (1.47 ± 0.06-fold; P = 0.002) and 10−8 M VIP (1.48 ± 0.18-fold; P = 0.012). VIP (10−8 M) treatment increased N-cadherin protein levels in bovine and human CE cells to 1.98 ± 0.28-fold (P = 0.005) and 1.17 ± 0.10 (range, 0.91–1.87)-fold (P = 0.050) of their respective controls. VIP antagonist (SN)VIPhyb diminished the VIP effect. VIP silencing resulted in deterioration of the hexagonal cell shape and decreased levels of VIP protein and mRNA, N-cadherin (but not connexin-43) mRNA and protein, and the antiapoptotic Bcl-2 protein.

CONCLUSIONS. Through its autocrine VIP, CE cells play an active role in maintaining the differentiated state and suppressing apoptosis in the corneal endothelium in situ. (Invest Ophtalmol Vis Sci. 2008;49:3491–3498) DOI:10.1167/iovs.07-1543

The corneal endothelium, a single-cell layer, functions to maintain the transparency of the cornea by acting as a barrier to the movement of fluid into the cornea and by actively pumping fluid out of the cornea. Although their developmental origin is the neural crest, the corneal endothelial (CE) cells express neuron-specific enolase and have limited regenerating capacity. The mechanisms through which the corneal endothelium maintains its differentiated state throughout life remain unknown. Elucidation of these mechanisms will increase the level of our understanding of the pathogenesis of CE decompensation.

In the corneal endothelium, N-cadherin is a differentiation marker; the expression of N-cadherin coincides with the formation of the CE cell layer during eye development.9,10 The calcium-dependent cell–cell junctional protein N-cadherin mediates cell adhesion through Ga2− dependent homophilic interaction of the extracellular domain and anchoring its cytoplasmic domain to the actin cytoskeleton.10 As such, N-cadherin plays important roles in shaping cells,11,12 in pattern formation in the developing retina,12 and in the dendritic spine morphogenesis of neurons.13 As an integral component of the cell–cell junctional complex, N-cadherin not only mediates intercellular adhesion strengthening,14,15 it functions as an adhesion-activated receptor capable of initiating distinctive signaling pathways16 leading to cell differentiation and survival. For example, N-cadherin adhesion induces differentiation and lamellipodium extension of myogenic cells,16,17 and stimulation of the N-cadherin signaling pathway increases expression of the antiapoptotic protein Bcl-218 and phosphorylation (inactivation) of the apoptotic protein BAD.19

The mechanisms that regulate N-cadherin gene expression remain largely unknown, though they have been shown to be CAMP dependent.20,21 In bovine and human CE cells, CAMP production is stimulated by vasoactive intestinal peptide (VIP),22 suggesting that VIP may play a role in the modulation of N-cadherin expression in CE cells. VIP is a 28-amino acid neuropeptide that binds to two types of adenylyl cyclase stimulatory heterotrimetric G protein–coupled receptors (VPAC1 and VPAC2) and transduces signal through CAMP-dependent and -independent pathways.23–25 While VIP is widely distributed in the central and peripheral nervous systems, including those in the eye,26,27 and in the immune system,28 we have previously reported that VIP mRNA and protein are also expressed in the CE cells in human and bovine corneas.29 Recently, we have reported that VIP expression in human CE cells in situ is upregulated by the injury factor ciliary neurotrophic factor (CNTF),29 which in turn is an autocrine released by CE cells surviving oxidative stress ex vivo.31 Furthermore, VIP is present and identified as one of the immunosuppressive factors in the aqueous humor,32–34 the fluid that fills the anterior and posterior chambers of the eye and that constantly bathes the CE cells in vivo. Therefore, through modulation of the CAMP level, the endogenous VIP in CE cells may play a role in regulating the N-cadherin level. Thus, in addition to being a trophic factor of CE cells under acute oxidative stress,29 VIP may also play a role as a differentiation-promoting factor by inducing N-cadherin synthesis in CE cells, in agreement with the role of VIP found in neuroendocrine differentiation.35,36 Neurite genesis and remodeling,37,38 and melanin genesis in retinal pigment epithelium.39

From the Departments of 1Ophthalmology and Visual Sciences and 2Physiology, University of Maryland, Baltimore, Maryland; and 3Transgenetech, Columbia, Maryland.

Supported in part by National Institutes of Health Grant RO1EY-11607 (S-WMK) and by Research to Prevent Blindness, Inc.

Submitted for publication December 3, 2007; revised February 5 and March 28, 2008; accepted June 19, 2008.

Disclosure: S.-W.M. Koh, None; K. Chandrasekara, None; C.J. Abbondandolo, None; T.J. Coll, None; A.R. Rutzen, None

The publication costs of this article were defrayed in part by page charge payment. This article must therefore be marked "advertisement" in accordance with 18 U.S.C. §1734 solely to indicate this fact.

Corresponding author: Shay-Whey M. Koh, Department of Ophthalmology and Visual Sciences, University of Maryland, Room 500C MSTF, 10 S. Pine Street, Baltimore, MD 21201; skoh@umaryland.edu.
While it is well known that the corneal endothelium in the adult human is a nonregenerating and differentiated tissue, the distinct possibility exists that the expression of the differentiation marker N-cadherin is maintained at a certain level, perhaps by VIP. In addition to testing the effect of exogenous VIP, the present study investigated effects of silencing endogenous VIP using lentiviral-based short hairpin RNA (shRNA) to knock down VIP mRNA, on the expression of N-cadherin and the normal hexagonal cell shape of CE cells in human corneas ex vivo. The present study is the first in silencing gene expression in CE cells in situ and in silencing VIP gene expression by RNA interference.

**Materials and Methods**

**Media**

**Medium A.** Medium A consisted of Dulbecco modified Eagle medium (DMEM) supplemented with 100 μg/ml penicillin, 100 μg/mL streptomycin sulfate, and 20 mM HEPES.

**Medium B.** Medium B consisted of DMEM supplemented with 100 μg/mL penicillin, 100 μg/mL streptomycin sulfate, and 0.292 mg/mL l-glutamine. Complete medium B consisted of medium B plus 5% fetal calf serum and fungizone (250 ng/mL amphotericin; Invitrogen; Grand Island, NY).

**Bovine Corneoscleral Explants**

Bovine eyes obtained from the local abattoir within 6 hours of kill were immersed in a 0.5% iodine solution (1:4 dilution in PBS of povidone iodine preparation solution [Baxter Health Care Co., Deerfield, IL]) for 5 seconds and rinsed in PBS before corneoscleral explants were dissected using a procedure similar to that described previously.40 A scleral incision 1 mm posterior to the limbus was applied, and the explant that was without the iris, ciliary body, or trabecular meshwork was lifted from the eyeball. Explants were rinsed once with 14 mL Dulbecco phosphate-buffered saline (DPBS) in 60 × 15-mm Petri dishes. With its concave side up, the explants were then incubated for 30 minutes at room temperature in 14 mL medium A in 60 × 15-mm Petri dishes and were transferred to 14 mL medium B in 60 × 15-mm Petri dishes and incubated at 37°C in 5% CO₂/95% air.

**Human Donor Corneas**

Viable human corneoscleral explants (human donor corneas) preserved for 5 to 21 days in storage medium (Optisol-GS; Chiron Ophthalmics, Irvine, CA) at 4°C were obtained from the Lions Eye Institute for Transplant & Research, Inc. (Tampa, FL). These corneas were deemed not suitable for transplantation because of the less than optimal CE cell densities and the advanced ages of the donors. Using the same procedure as the eye bank, fresh human donor corneoscleral explants were retrieved from cadavers (within 30 hours after death) in the Anatomy Board of the State of Maryland. The cadavers have been identified and were not considered human subjects by the Human Research Protection Office, University of Maryland School of Medicine. Human corneoscleral explants were established in 24-hour organ cultures (with 3.5 mL medium) as the bovine corneoscleral explants.

**VIP Treatment of Bovine Corneoscleral Explants and Human Donor Corneas**

**Bovine Corneoscleral Explants.** Twenty-four hours after their establishment in organ cultures, bovine explants were transferred to and incubated in 35-mm culture dishes containing 4 mL fresh medium B plus 5% fetal calf serum and VIP (0, 10⁻¹² to 10⁻⁸ M) for 25 to 48 hours at 37°C in 5% CO₂/95% air. To test the effects of blocking VIP binding to its receptor, VIP antagonist (SNV)Phyb (Bachem, Torrance, CA) was included in the medium during the VIP treatment.

**Human Donor Corneas.** Paired human donor corneas were removed from the storage medium (Optisol-GS; Bausch & Lomb, Irvine, CA)-containing storage vials and incubated in 2 mL complete medium B and either zero (OD) or 10⁻⁸ M VIP (OS) in 24-well plates for 48 hours at 37°C in 5% CO₂/95% air.

**Lentiviral Transduction of Paired Fresh Human Donor Corneas**

Various lentiviral vectors have been shown, including in human in vitro and vivo studies,41-44 to efficiently and persistently transduce genes into CE cells of various species. In the present study, conditions of transduction were first characterized closely according to the protocol provided by the manufacturer of the lentiviral particles used (Mission TurboGFP; Sigma-Aldrich, St. Louis, MO).

In the initial experiments to determine the efficiency of lentiviral transduction, lentiviral particles that express green fluorescent protein (GFP, Mission TurboGFP control transduction particles SHC003V; Sigma-Aldrich) were added in increasing amounts to 90% confluent SK-N-SH neuroblastoma cells (American Type Culture Collection, Manassas, VA) seeded at 1.6 × 10⁵ cells/dish in complete medium B in 35-mm culture dishes. To enhance lentiviral transduction, hexadimethrine bromide was added to the culture medium at a final concentration of 8 μg/mL. After 24 hours of exposure to lentiviral particles, the number of GFP-positive SK-N-SH cells was determined. The optimal amount of lentiviral particles (15 μL in 1 mL complete medium B) that resulted in GFP expression in 50% of SK-N-SH cells was determined and used to transduce CE cells in fresh human donor corneas. A representative photomicrograph of a flat-mounted human donor cornea (Supplementary Fig. S1, online at http://www.iovs.org/cgi/content/full/49/8/3491/DC1), showing CE cells transduced with GFP-lentiviral particles (in the presence of 0.5 μg/mL hexadimethrine bromide) for 72 hours, expressed green fluorescent GFP.

The next set of experiments screened five distinctive lentiviral particles produced from five short hairpin RNA (shRNA) lentiviral plasmid vectors. Each could generate intracellularly a distinctive short interfering RNA (siRNA) targeting one of the five human VIP gene sequences (TRCN0000078053-TRCN0000078057; Mission Lentiviral Transduction Particles, SHVRS, Sigma-Aldrich) for the most effective lentiviral particle that caused maximal decrease in VIP mRNA. SK-N-SH cells were exposed to lentiviral particles for 24 hours, as described, the cells were rinsed free of viral particles with DPBS, total RNA was isolated, and semiquantitative RT-PCR for VIP was performed. The lentiviral particle that showed a maximal decrease in VIP mRNA expression was determined to be the shRNA with the following sequence: CCGGCCCGACTTTCCAGAAGAGTTACTCGAGTAACTCTTCT- GAAAGTCGGGTTTTTG (TRCN0000078057), targeting VIP mRNA sequences 656-676 (NM003381.2) or 653–673 (NM 194435.1).

To study the effect of VIP gene silencing in CE cells, paired fresh human corneas from a 97-year-old female (donor 1), an 81-year-old female (donor 2), a 76-year-old female (donor 3), a 69-year-old female (donor 4), and a 70-year-old female (donor 5) were used in four separate experiments. Within each pair, one of the corneas was transduced with lentiviral particles (TRCN0000078057) determined to be most efficient in silencing VIP gene expression, whereas the paired cornea was transduced with the negative control lentiviral particles produced from the pLKO.1-puro control vector (SHC001V; Sigma-Aldrich). Toward this end, 10 μL stock solutions containing 10⁵ to 10⁶ lentiviral particles in DMEM with 10% heat-inactivated fetal bovine serum and antibiotics was added to the 0.5 mL complete B medium (plus 0.5 μg/mL hexadimethrine bromide) incubating the corneas. After 72 hours, corneas of donors 1, 2, and 5 were bisectioned. Half the cornea was used to isolate CE cell total RNA for semiquantitative RT-PCR, and the other half was further bisected. One quadrant was stained with alizarin red S in calcium and magnesium-free DPBS, followed by washing 2× with DPBS and examination under a microscope, and the other quadrant was fixed with 4% paraformaldehyde in PBS for VIP immunostaining. From each cornea of donors 4 and 5, a quadrant was dissected, stained with alizarin red S, washed 2× with DPBS, and examined under a microscope. From the remaining corneas, CE cells were scraped from the Descemet membrane and extracted for West-
ern blot analysis. A second lentiviral particle that was not as effective as TRCN0000078057 in knocking down VIP mRNA expression in transduced CE cells was used to corroborate the results obtained with TRCN0000078057. This second lentiviral particle (TRCN0000078053) has the shRNA with the sequence CCGGGCTGTGTTAAATAAACCTT/CATACTCACAATGAGAC-3′ [forward] and 5′-CTGAGAACTGATAAGAAAAATCGT-3′ [reverse].45,46 (2) human connexin 43 (5′-CCCTTCTGGATGCAGTGTAC-3′ [forward] and 5′-ACCAAGGACCCACACAGCAT-3′ [reverse]),47 and (3) human VIP (5′-CCTGTCTTATGGGAAAACGTGT-3′ [forward] and 5′-H11032 [reverse]),48,49 (2) human connexin 43, (3) human VIP, and (4) human N-cadherin (BTA7 [R&D Systems, Minneapolis, MN] 205606 [Calbiochem, San Diego, CA]). The difference of the control versus VIP-treated corneoscleral explants was significant at the levels of *P < 0.05, **P < 0.01, and ***P < 0.002.

Western Blot Analysis

Samples of CE cell extracts were electrophoresed under reducing conditions using preform Tris/Glycine polyacrylamide gradient gels (NuPage; Novex, San Diego, CA) and electrophoretically transferred to nitrocellulose membranes for Western blot analysis using the following primary antibodies: affinity-purified polyclonal antibodies against N-cadherin (BTA7 [R&D Systems, Minneapolis, MN]) anti–VIP rabbit serum (T-4246 [Peninsula Laboratories, San Carlos, CA]), and mouse monoclonal antibody against Bcl-2 (05–729 [Upstate Biotechnology, Lake Placid, NY]). For VIP, a secondary antibody–biotinylated goat anti–rabbit IgG in conjunction with extravidin–alkaline phosphatase conjugate (E-2636; Sigma) and a kit (Fast Red TR/Naphthol AS-MX tablet set; Sigma) were used to develop the red reaction product. To reprobe the blots with a monoclonal actin antibody (Ab-1 kit; Oncogene, Cambridge, MA), the blots were stripped with a buffer (Restore Western Blot Stripping buffer; Pierce, Rockford, IL) according to the instructions of the manufacturer. Densities of immunoreactive bands were determined same as in semiquantitative RT-PCR.

VIP Immunostaining

After lentiviral transduction quadrants of human donor corneas were fixed in 4% paraformaldehyde in PBS, embedded in paraffin, and sectioned (6 μm) for immunostaining using an anti–VIP rabbit serum (T-4246; 1:1000 in PBS; Peninsula Laboratories, San Carlos, CA) and a secondary antibody biotinylated goat anti–rabbit IgG. Extravidin–alkaline phosphatase conjugate (E-2636; Sigma) and a kit (Fast Red TR/Naphthol AS-MX tablet set; Sigma) were used to develop the red reaction product in positively stained cells. Rabbit IgG (10 μg/mL) was used as a negative control.

CE Cell Extract

**Bovine CE Cell.** At the conclusion of VIP treatment of the corneoscleral explants, corneoscleral explants were removed from the medium, and the corneal endothelium was scraped off the corneas using a razor blade and homogenized in the RIPA buffer, as described.

**Human Corneas Previously Stored in Medium.** The endothelium and Descemet membrane were hydrosedimented from the posterior stroma of human corneoscleral explants using the same method previously described (Rutzen AR, et al. IOVS 2000;41:ARVO Abstract 2376). An injection of DPBS at the posterior limbus was used to create a fluid separation between the endothelium/Descemet membrane and posterior stroma. A small incision was created at the peripheral posterior limbus to release the DPBS from beneath Descemet membrane. A 7.5-mm trephine was used to excise the specimen of endothelium/Descemet membrane, which was then treated as described for the bovine corneal endothelium.

**Human Corneas from Cadavers.** The corneal endothelium was scraped from each cornea using a razor blade and homogenized in the RIPA buffer, as described.

**Semiquantitative RT-PCR**

Total RNA was isolated from CE cells scraped from bovine and lentiviral particles–transduced human corneas (RNA-Be; Tel-Text, Friendswood, TX) and subjected to reverse transcription (RT) using a kit (RETrOscrip; catalog number 1710; Ambion; Austin, TX). RT products were subjected to PCR using three primer sets corresponding to the gene sequences of (1) human N-cadherin (5′-CACCACACATGTTTACAATCAAAATGAGAC-3′ [forward] and 5′-CTGAGAACTGATAAGAAAAATCGT-3′ [reverse]),45,46 (2) human connexin 43 (5′-CCCTTCTGGATGCAGTGTAC-3′ [forward] and 5′-ACCAAGGACCCACACAGCAT-3′ [reverse]),47 and (3) human VIP (5′-CCTGTCTTATGGGAAAACGTGT-3′ [forward] and 5′-H11032 [reverse]).48,49 For quantifying mRNA levels, semiquantitative PCR was conducted in the presence of the 18S primers and the 18S competomers (Universal 18S internal standards; catalog number 1718; Ambion) at a ratio of either 1:9 (N-cadherin and connexin 43) or 0:5:9:5 (VIP). PCR used RT products derived from 0.15 μg bovine RNA (for N-cadherin), 10% of isolated human RNA (for N-cadherin and connexin 43), and 20% of isolated human RNA (for VIP). Between an initial 2-minute (94°C) and a final 5-minute (72°C) treatment, 25 to 38 thermocycles (20 seconds at 94°C, 25 seconds at the annealing temperature, and 40 seconds at 72°C) were conducted. The numbers of cycling and the annealing temperatures (in °C) were proven by sequencing (Biopolymer Laboratory, University of Maryland) and were demonstrated to be identical with the sequences for bovine N-cadherin (442 bp), human N-cadherin (444 bp), human connexin 43 (154 bp), and human VIP (86 bp).

**Figure 1.** CE cell N-cadherin mRNA level increased in a VIP-concentration dependent manner. (A) N-cadherin and 18S mRNA levels reflected in the electrophoresed (2% agarose) semiquantitative RT-PCR products. (B) The ratio of normalized CE cell N-cadherin cDNA (against the 18S internal standard) levels over that averaged from control corneoscleral explants of the same experiment (y-axis) was shown as a function of VIP concentration. Data were combined from three separate experiments. The difference was significant (P < 0.05, ANOVA) among various groups: corneoscleral explants treated with 0 N = 6, 10−12 (N = 3), 10−10 (N = 3), 10−8 (N = 3), and 10−6 (N = 5) M VIP. The difference of the control versus VIP-treated corneoscleral explants was significant at the levels of *P < 0.02, **P < 0.01, and ***P < 0.002.
**RESULTS**

**VIP-Concentration Dependency of N-Cadherin mRNA Levels in CE Cells in VIP-Treated Corneoscleral Explants**

To determine the maximally effective VIP concentration in increasing the N-cadherin level, bovine corneoscleral explants were used. VIP (0, 10^{-12}, 10^{-10}, 10^{-8}, and 10^{-6} M) treatment for 25 hours resulted in increased N-cadherin mRNA levels in CE cells in a VIP concentration-dependent manner (Fig. 1). Semiquantitative RTPCR of N-cadherin and 18S (internal standard) mRNAs showed that the N-cadherin levels in CE cells in 10^{-12}, 10^{-10}, 10^{-8}, and 10^{-6} M VIP-treated corneoscleral explants increased (in mean ± SEM) to 1.23 ± 0.07-fold (P < 0.01), 1.47 ± 0.14-fold (P < 0.002), 1.48 ± 0.24-fold (P < 0.02), and 1.41 ± 0.14-fold (P < 0.01) of the control corneoscleral explants (significance levels of VIP-treated versus control are shown in parentheses). The difference among various groups was significant at P < 0.05 (ANOVA) (Fig. 1B).

**Increased N-Cadherin Protein Level in CE Cells in VIP-Treated Bovine Corneoscleral Explants and Human Donor Corneas**

To show the effect of VIP on the protein level of CE cell N-cadherin, bovine corneoscleral explants and paired human donor corneas were treated with VIP at the concentration that produced the maximal effect on increasing the N-cadherin mRNA level (Fig. 1).

After 48-hour treatment of bovine corneoscleral explants with 10^{-8} M VIP, CE cells were analyzed by Western blot analysis for N-cadherin and actin (as internal standard). As shown in Figure 2A, the N-cadherin level in CE cells from VIP-treated corneoscleral explants was higher than that in the control. Figure 2B showed that VIP treatment increased N-cadherin levels to 1.98 ± 0.28-fold (mean ± SEM; N = 8) of that found in the control corneoscleral explants (N = 7, P = 0.005).

The effect of VIP (1 × 10^{-8} M VIP) was diminished by the presence of the VIP antagonist (SN)VIPhyb (5 × 10^{-7} M) during the VIP treatment of the bovine corneoscleral explants (Figs. 2C, 2D). The level of connexin 43, a protein localized to the gap junction, was not affected by the VIP treatment (data not shown).

**Knockdown of Endogenous VIP Expression in Human CE Cells by Lentiviral Based shRNA Leading to Deterioration of the Hexagonal Cell Shape and Decreased N-Cadherin Level**

Given that VIP protein and mRNA are expressed by CE cells, knocking down the endogenous VIP expression would lead to disturbances of the VIP signaling pathways and would help identify the normal functions of endogenous VIP. Toward this end, three pairs of fresh human donor corneas were used. In each pair, one cornea was transduced with shRNA lentiviral particles targeted at knocking down VIP gene expression, whereas the paired cornea was transduced with the negative control versus VIP (10^{-8} M)-treated. Western blot analysis of CE cells showed that VIP was effective in increasing the N-cadherin level (Fig. 3A) in six of nine human donor cornea pairs. In all nine pairs, VIP-treated corneas demonstrated N-cadherin levels that were 0.91- to 1.87-fold of their respective controls (1.173 ± 0.099; P = 0.050; Fig. 3B). There was no correlation between VIP responsiveness and either donor age or cause of death (data not shown).

**FIGURE 2.** VIP (10^{-8} M) treatment of bovine corneoscleral explants increased the protein level of N-cadherin in CE cells (A, B), and VIP antagonist (SN)VIPhyb (5 × 10^{-7} M) diminished the VIP effect (C, D). (A) Western blot depicting N-cadherin and actin-immunoreactive bands. Each lane contained 32 μg protein. (B) The ratio of normalized N-cadherin level (against the actin internal standard) over the average of those found in the control corneoscleral explants of the same experiment. Data were combined from five separate experiments showing VIP treatment of corneoscleral explants (N = 8) increased the CE cell N-cadherin level to 1.98 ± 0.28-fold of the control (N = 7) (P = 0.0048). (C) Western blot showing the effect of VIP antagonist on diminishing the CE cell N-cadherin level in VIP-treated corneoscleral explants. (D) Normalized N-cadherin levels (against the actin internal standard) in VIP-treated corneoscleral explants in the absence and presence of the VIP antagonist averaged 2.27 ± 0.33 (N = 4) and 1.09 ± 0.13 (N = 3), respectively (P = 0.017). The data were representative of three separate experiments with similar results.

**FIGURE 3.** VIP-increased N-cadherin levels demonstrated in paired donor human corneas, as control versus VIP (10^{-8} M) treated. (A) Western blot analysis of N-cadherin and actin (as internal standard) in CE cells of control (Con) and VIP-treated (VIP) paired corneas. Lanes contained extract of CE cells in sheets of corneal endothelium (7.5 mm in diameter) trephined from the centers of the corneas. (B) Normalized N-cadherin level (against the actin internal control) of nine pairs of human corneas, ranked by the effectiveness of VIP, from the lowest (donor 1) to the highest (donor 9). VIP treatment increased N-cadherin level to (mean ± SEM) 1.173 ± 0.10-fold of the control (P = 0.050). Ages of donors 1 to 9 were, respectively, 75, 58, 72, 64, 66, 73, 61, 71, and 67 years.
control lentiviral particles. VIP gene knockdown dramatically affect the morphology of the CE mosaic (Figs. 4A, 4B). Alizarin red S staining of the cell–cell junctions revealed in flat-mounted corneas the normal hexagonal CE cell shape in the control (A) and deterioration of the hexagonal cell shape after VIP knockdown (B). Downregulation of CE cell VIP expression was demonstrated by VIP immunostaining of 6-μm corneal sections (C, D), semiquantitative RT-PCR followed by agarose gel electrophoresis of RT-PCR products (E), densitometry of the VIP and 18S (internal standard) bands in the gel (F), and Western blot analysis (G) followed by densitometry (H) of the VIP-immunoreactive molecule and the Ponceau S–stained 65-kDa molecule, which was the most abundant protein in CE cell extract. Identical results were obtained from all five pairs (A, B) and both donor pairs (donors 1 and donor 2) (C–F) examined. (F) Data were averaged from the two pairs; the third pair did not yield a sufficient number of CE cells for RT-PCR. Two additional pairs of corneas (from donors 4 and 5) were used in Western blot analysis for the CE cell VIP immunoreactive molecule (G, H). Original magnifications: ×200 (A, B); ×1000 (C, D).

FIGURE 4. Lentiviral-mediated VIP shRNA in human CE cells altered cell shape and decreased VIP expression. Five pairs of human donor corneas were used. Within each pair, one cornea was transduced with control lentiviral particles (Con) and the other cornea was transduced with those with the VIP mRNA-interfering nucleotide sequence (Kno). Alizarin red S staining of the cell–cell junctions revealed in flat-mounted corneas the normal hexagonal CE cell shape in the control (A) and deterioration of the hexagonal cell shape after VIP knockdown (B). Downregulation of CE cell VIP expression was demonstrated by VIP immunostaining of 6-μm corneal sections (C, D), semiquantitative RT-PCR followed by agarose gel electrophoresis of RT-PCR products (E), densitometry of the VIP and 18S (internal standard) bands in the gel (F), and Western blot analysis (G) followed by densitometry (H) of the VIP-immunoreactive molecule and the Ponceau S–stained 65-kDa molecule, which was the most abundant protein in CE cell extract. Identical results were obtained from all five pairs (A, B) and both donor pairs (donors 1 and donor 2) (C–F) examined. (F) Data were averaged from the two pairs; the third pair did not yield a sufficient number of CE cells for RT-PCR. Two additional pairs of corneas (from donors 4 and 5) were used in Western blot analysis for the CE cell VIP immunoreactive molecule (G, H). Original magnifications: ×200 (A, B); ×1000 (C, D).

control lentiviral particles. VIP gene knockdown dramatically affect the morphology of the CE mosaic (Figs. 4A, 4B). Alizarin red S staining of the cell–cell borders in the corneal quadrant from all three pairs of corneas (see Materials and Methods) demonstrated that VIP gene knockdown resulted in deterioration of the hexagonal cell shape and appearance of irregularly shaped CE cells. Each individual CE cell in VIP knockdown corneas covered a larger area than that in the control corneas (Figs. 4A, 4B). Immunocytochemical studies of the VIP protein expression (Figs. 4C, 4D), semiquantitative RT-PCR of VIP mRNA (Figs. 4E, 4F), and Western blot analysis (Figs. 4G, 4H) confirmed the effectiveness of the VIP shRNA lentiviral particles in knocking down endogenous VIP expression. The normalized VIP (against 18S) mRNA level in the control CE cells of two human corneas (donors 1 and 2) averaged 0.49, whereas that in the VIP knockdown CE cells in the paired human corneas was 0.28, a 43% reduction. Previously, a specific VIP-immunoreactive molecule with the same molecular size as the prepro-VIP (20 kDa) has been demonstrated in fresh human CE cells.29 In two pairs of human corneas (donors 4 and 5), VIP gene knockdown in CE cells reduced the level of the 20-kDa VIP-immunoreactive molecule (Fig. 4G) to 18% of that found in control corneas (Fig. 4H).

Deterioration of the hexagonal CE cell shape was accompanied by a decrease in N-cadherin expression in VIP knockdown CE cells. As shown in Figure 5A, semiquantitative RT-PCR of N-cadherin mRNA demonstrated that VIP knockdown in CE cells of human corneas decreased the N-cadherin mRNA level.
The normalized N-cadherin (against 18S) mRNA levels in the control CE cells of two human corneas (from two donors) and those in the VIP knockdown CE cells of the paired corneas averaged 10.2 and 2.3, respectively (Fig. 5B). In the meantime, VIP gene expression knockdown did not affect the expression of connexin 43, a gap junctional protein (Figs. 5C, 5D).

To show the effect of VIP gene knockdown on the protein level of CE cell N-cadherin, Western blot analysis was performed using four pairs of human corneas. Figure 6A demonstrates results from two pairs of corneas (donors 4 and 5). In VIP gene knockdown CE cells transduced by TRCN0000078057 particles, the N-cadherin protein level was drastically reduced; the protein levels of actin and Bcl-2 were also reduced, but to lesser extents. In VIP gene knockdown CE cells, the protein levels of N-cadherin, actin, and Bcl-2 were 1% (Fig. 6B), 31% (Fig. 6C), and 54% (Fig. 6D) of their respective controls. Results were corroborated with those from two additional pairs of corneas (donors 6 and 7), in which CE cells were transduced with the second lentiviral particle (TRCN0000078053), which was not as effective as the TRCN0000078057 particle in knocking down VIP gene expression (see Materials and Methods), resulting in decreases in the protein levels of N-cadherin, actin, and Bcl-2 to 11% (Fig. 6B), 67% (Fig. 6C), and 90% (Fig. 6D) of their respective controls.

**DISCUSSION**

The present study demonstrated that the CE cell autocrine VIP increased N-cadherin expression and maintained the normal hexagonal cell shape in the corneal endothelium, whereas our previous reports demonstrate that VIP protects the corneal...
endothelium to the acute killing effect of oxidative stress,²⁹ that endogenous VIP is upregulated by CNTF,³⁰ and that CNTF in turn is released by CE cells surviving oxidative stress.³¹ We therefore hypothesize that the CE cell autocrine factors CNTF and VIP work in concert to maintain the differentiated state and to promote the survival of the corneal endothelium.

Although a few growth factors such as the hepatocyte growth factor and epidermal growth factor have been shown to upregulate N-cadherin expression,⁴⁶ how N-cadherin gene expression is regulated is unknown. The proximal promoter sequence of the N-cadherin gene has been characterized.⁴⁸ It has high GC content and contains an Sp1/Sp3-binding site, which functions as the basal regulatory element of the promoter. Whereas Sp1 binding to the promoter is increased by cAMP,⁴⁹ activated CAMP signaling pathways increase the activity of the Sp1-binding site-containing promoter of various genes.⁵⁰,⁵¹ In particular, the Sp1-binding site is required for cAMP-dependent, follicle-stimulating hormone (FSH)-stimulated immature-like growth factor binding protein gene expression.⁵² Although FSH-stimulated N-cadherin gene expression in Sertoli cells is mediated by cAMP,²⁰ the role of the Sp1-binding site in camp-stimulated N-cadherin gene expression has not been demonstrated. We have previously demonstrated VIP-stimulated cAMP production in CE cells in human and bovine corneas.²² The observation in the present study—supported by our preliminary study showing that actinomycin dose dependently diminished the effect of VIP on increasing N-cadherin level—of VIP-increased levels of N-cadherin was likely the result of VIP upregulation of N-cadherin gene expression.

Various lentiviral vectors efficiently and persistently transduce genes into CE cells of several species, including humans, as observed in vitro and in vivo studies, resulting in no detrimental effect on the corneal endothelium morphology.⁴¹–⁴⁴ What is the sequence of events leading to the deterioration of the normal hexagonal geometry of the apical surface of the CE cells after VIP gene knockdown (Figs. 4A, 4B)? VIP knockdown decreased only N-cadherin, but not connexin 43, expression (Fig. 5), and exogenous VIP increased N-cadherin level in CE cells (Figs. 1–3) but had no effect on that of connexin 43 (data not shown). Furthermore, loss of N-cadherin, but not of E-cadherin, alters cone cell shape in Drosophila eyes.³¹ We tentatively concluded that, in the VIP knockdown CE cells, decreased VIP expression (Figs. 4C–H) led to decreased levels of N-cadherin (Figs. 5A, 5B, 6A, 6B), cell–cell adhesion, actin (Figs. 6A, 6C), and stability of the actin cytoskeleton to which the cytoplasmic domain of N-cadherin normally anchors,¹⁰ and to the deterioration of the normal hexagonal cell shape. Future studies will investigate the possibility that exogenous VIP may rescue the VIP knockdown CE cell and restore its hexagonal shape. Future studies of N-cadherin gene knockdown will ascertain the cause-and-effect relationship between the loss of N-cadherin expression and hexagonal cell shape.

After VIP knockdown, the area covered by individual CE cells increased (Figs. 4A, 4B), indicative of cell spreading, which is a well-recognized repair mechanism in this nonregenerating tissue. Consistent with this finding was that more denuded areas devoid of CE cells were present in the VIP knockdown corneas (Figs. 4A, 4B). Given that VIP is a trophic factor capable of protecting CE cells against oxidative stress,⁴⁹ reduced endogenous VIP levels might have increased the vulnerability of CE cells to stress induced by lentiviral transduction.³⁴ In addition, the signaling pathway that is initiated by N-cadherin in the junctional complex and that leads to increased levels of antiapoptotic protein Bcl-2 and phosphorylation (inactivation) of the proapoptotic protein BAD might have been affected as a result of VIP gene knockdown.

Indeed, decreased levels of the antiapoptotic protein Bcl-2 (Figs. 6A, 6D) were observed in CE cells after the VIP gene was knocked down by either the most effective lentiviral particles (TRCN0000078057) or the less effective (see Materials and Methods) ones (TRCN0000078053). Although the extent of Bcl-2 reduction correlated well with that of N-cadherin such that the more N-cadherin was downregulated the more Bcl-2 level was reduced (Figs. 6A, 6B, 6D), N-cadherin expression was more sensitive to VIP gene knockdown than Bcl-2, suggesting the involvement of other signaling pathways in the regulation of Bcl-2 expression. In the present study, phosphorylated BAD was not detected in the Western blot analysis of CE cells either in the control corneas or in those with VIP gene knockdown. It remains to be studied whether the level of BAD was higher in VIP knockdown CE cells; these results nonetheless predicted increased apoptotic CE cell death after VIP gene knockdown. Furthermore, because N-cadherin mediates intercellular adhesion strengthening,¹³,¹⁴ CE cells with decreased N-cadherin levels resulting from VIP knockdown may detach from the corneal endothelium sheet.

In conclusion, the present study demonstrated for the first time a definite role of an autocrine in maintaining the differentiated state of the corneal endothelium and the feasibility of using RNA interference for studying the functions of specific genes in corneal endothelium in situ.

Acknowledgments

The authors thank the Lions Eye Institute for Transplant & Research, Inc. (Tampa, FL) and the Anatomy Board of the State of Maryland (Baltimore, MD) for the human donor corneas, and Dante Gloria and Jason M. Cheng for excellent technical support.

References


Downloaded From: https://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/932954/ on 11/26/2018
3498 Koh et al.


