MicroRNA-184 Regulates Corneal Lymphangiogenesis

Sammy Grimaldo, Don Yuen, Jaci Theis, Melissa Ng, Tatiana Ecoiffier, and Lu Chen

Center for Eye Disease and Development, Program in Vision Science, and School of Optometry, University of California, Berkeley, California, United States

PURPOSE. MicroRNAs are a class of small noncoding RNAs that negatively regulate gene expression by binding to complimentary sequences of target messenger RNA. Their roles in corneal lymphangiogenesis are largely unknown. This study was to investigate the specific role of microRNA-184 (mir-184) in corneal lymphangiogenesis (LG) in vivo and lymphatic endothelial cells (LECs) in vitro.

METHODS. Standard murine suture placement model was used to study the expression change of mir-184 in corneal inflammatory LG and the effect of synthetic mir-184 mimic on this process. Additionally, a human LEC culture system was used to assess the effect of mir-184 overexpression on cell functions in vitro.

RESULTS. Expression of mir-184 was significantly downregulated in corneal LG and, accordingly, its synthetic mimic suppressed corneal lymphatic growth in vivo. Furthermore, mir-184 overexpression in LECs inhibited their functions of adhesion, migration, and tube formation in vitro.

CONCLUSIONS. These novel findings indicate that mir-184 is involved critically in LG and potentially could be used as an inhibitor of the process. Further investigation holds the promise for divulging new therapies for LG disorders, which occur inside and outside the eye.

Keywords: lymphangiogenesis, cornea, microRNA, lymphatic endothelial cell

Moreover, mir-184 overexpression in human lymphatic endothelial cells (LECs) in vitro suppresses their functions of adhesion, migration, and tube formation. These results together reveal that mir-184 is a negative regulator of the lymphangiogenic process. Further investigation on this natural regulator holds the great promise for developing new and effective treatment for LG-related diseases in the body.

METHODS

Animals, LECs, and Reagents

Normal adult 6- to 8-week-old male BALB/c mice were purchased from Taconic Farms (Germantown, NY, USA). All mice were treated according to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research and the protocols approved by the Animal Care and Use Committee of the institute. Mice were anesthetized using a mixture of ketamine, xylazine, and acepromazine (50, 10, and 1 mg/kg body weight, respectively) for each surgical procedure. Human neonatal primary microdermal LECs were purchased from Lonza (Walkersville, MD, USA) and maintained in EGM-2MV cell culture medium (Lonza) according to manufacturer's instructions. Matrigel, collagen type I, and calcine AM were purchased from BD Biosciences (San Jose, CA, USA). Mir-184 mimic and control RNA were purchased from Dharmacon, Inc. (Lafayette, CO, USA) and Ambion (Austin, TX, USA).

Induction of Corneal LG and Pharmaceutical Intervention

The experiments were performed as described previously. A standard suture placement model was used to induce corneal...
Digital images were taken with an epifluorescence microscope (Axiolmager M1; Carl Zeiss AG, Göttingen, Germany) and analyzed using the National Institutes of Health (NIH; Bethesda, MD, USA) ImageJ software (available in the public domain at http://imagej.nih.gov/ij/). The percentage scores of LG coverage areas were obtained by normalizing to control groups defined as being 100%. The differences were analyzed using the Mann-Whitney U test with P < 0.05 as significant.

Reverse Transcription and Real-Time PCR
Total RNA from LECs or central corneal epithelium was extracted using miRNeasy Mini Kit (Qiagen, Valencia, CA, USA). Reverse transcription was performed using the miScript II RT Kit (Qiagen). Real-time PCR was performed using miScript SYBR Green PCR Kit with specific primers to mature mir-184 using the mir-184_1 miScript Primer Assay (Qiagen) and measured by the CFX96 real-time detection system (Bio-Rad, Hercules, CA, USA). Relative expression of the mir-184 was calculated from the \( \Delta\Delta C_{T} \) (threshold cycle) of the targeted gene normalized to the \( \Delta\Delta C_{T} \) of the RNU6B reference gene. Tradional PCR products also were run on a 2% agarose gel.

Mir-184 Ectopic Expression in LECs
Transfections were done with Lipofectamine RNAiMax (Invitrogen, Carlsbad, CA, USA) according to manufacturer’s instructions. As reported previously, the transfection of either mir-184 mimic or control RNA was done overnight at 37°C in a 5% CO₂ humidified air incubator.

Adhesion Assay
The experiment was performed as described previously. At 72 hours following the transfection, 100 μL cells \( (3 \times 10^5 \text{ cells/mL}) \) were added to collagen type I-coated 96 plate wells and incubated for 30 minutes at 37°C. The plates then were washed several times and incubated with calcein (1 μg/mL) in Hank’s buffered salt solution (HBSS) for 30 minutes at room temperature. Plates were washed with PBS and fluorescent intensity from bound cells was measured with a Spectramax M5e microplate reader (Molecular Devices, Sunnyvale, CA, USA). Assays were performed in triplicate and repeated at least three times.

Migration Assay
At 72 hours following the transfection, a 200-μL pipette tip was used to create linear wounds within LEC monolayers. Phase images of the scratches were taken at time 0 and 29 hours using a Zeiss Axio Observer A1 inverted microscope (Carl Zeiss AG). For better visualization of the scratch area by end of the study at 29 hours, the cells were stained with crystal violet. The TScratch program (Tobias Gebäck and Martin Michael Peter Schulz, Swiss Federal Institute of Technology, ETH Zurich, Zurich, Switzerland) was used to determine the percent of the open area.

Tube Formation Assay
As previously reported, 72 hours following transfection, LECs were seeded \( (2 \times 10^5 \text{ cells/well}) \) onto 96-well plates containing solidified Matrigel and monitored for 24 hours under a Zeiss Axio Observer A1 inverted microscope (Carl Zeiss AG). Phase images of tubes were taken and total tube
lengths were analyzed by NIH ImageJ software. Assays were performed in triplicate and repeated at least three times.

**Statistical Analysis**

The mean difference was analyzed by Student’s *t*-test using Prism software (GraphPad, La Jolla, CA, USA) unless otherwise indicated. The differences between the treatment and control groups were considered statistically significant when *P* < 0.05.

**RESULTS**

**Mir-184 is Downregulated in Corneal LG**

To investigate the role of mir-184 in corneal inflammatory LG, we first assessed the expression change of mir-184 in the inflamed cornea following suture placement and lymphatic ingrowth. As shown in Figure 1A by real-time PCR analysis, the expression level of mir-184 in the inflamed cornea was significantly downregulated, compared to the normal condi-
MicroRNA-184 Regulates Corneal Lymphangiogenesis

IOVS | November 2015 | Vol. 56 | No. 12 | 7212

Mir-184 Overexpression Inhibits LEC Adhesion

Adhesion is an important function of LECs in the lymphangiogenic process. We next determined whether mir-184 regulates LEC adhesion in vitro. At 72 hours following the transfection with mir-184 mimic or control RNA, LECs were subjected to a collagen 1 adhesion assay, as reported previously. Our results showed that mir-184 overexpression in LECs led to a significant reduction in cell adhesion (Fig. 3A, \( P < 0.05 \)), suggesting a negative inhibitory role of mir-184 in this function.

Mir-184 Overexpression Reduces LEC Migration

To assess whether mir-184 also is involved in LEC migration in vitro, we performed the wound healing scratch assay 72 hours following the transfection with mir-184 mimic or control RNA. As presented in Figures 3B and 3C, mir-184 transfected cells showed a significant decrease in the rate of wound closing with larger open area (\( P < 0.05 \)). These results indicated that mir-184 negatively regulates LEC migration as well.

Mir-184 Overexpression Suppresses LEC Tube Formation

We also examined the effect of mir-184 overexpression on the ability of LECs to organize into capillary-type tubes using a three-dimensional (3D) culture system. At 72 hours following the transfection with either mir-184 mimic or control RNA, LECs were seeded on Matrigel, a basement membrane matrix, and observed for 24 hours. As shown in Figures 4A and 4B, mir-184 ectopic expression revealed a significant reduction in total tube length (\( P < 0.05 \)), confirming an inhibitory role of mir-184 in this important LEC function in vitro.

DISCUSSION

In summary, to our knowledge we provided the first evidence that mir-184 negatively regulates the LG process. We reported two important findings: Mir-184 expression is significantly downregulated in corneal inflammatory LG, and mir-184 mimic can be used to suppress corneal LG in vivo and LEC functions in vitro. Taken together, this study not only divulges mir-184 as a natural suppressor of LG, but also puts forth the use of mir-184 mimics as a novel strategy for LG therapy.

Mir-184 has a restrictive expression profile in the cornea, brain, and testes. Previously reported microarray analysis revealed that it is one of the most abundantly expressed microRNAs in the corneal epithelium. In the current study, we showed that mir-184 is significantly downregulated during corneal inflammatory LG, and it is highly indicated that this microRNA may act to maintain the alymphatic status of the cornea under normal condition. Allied to this speculation is our additional data showing that reintroduction of mir-184 into the cornea suppresses lymphatic formation. As an immune privileged tissue, the cornea has developed mechanisms that actively maintain its lymphatic-free status under normal condition, which are yet to be fully explored and understood.

Recently, it was reported that a soluble VEGFR-2 is secreted by the corneal epithelium and acts to suppress LG. This current study not only divulges mir-184 as a new and natural inhibitor of LG, but also indicates that multiple mechanisms and factors are involved in maintaining the alymphatic status of the cornea, which warrants further investigation.

It is known that mir-184 mutation results in corneal pathologies in keratoconus and the endothelial dystrophy, iris hypoplasia, congenital cataract and stromal thinning (EDICT) syndrome. To date, there has been no study linking mir-184 to corneal pathologic LG. Previously, it was reported that mir-184 is involved in ischemia-induced retinal angiogenesis, and the downstream targets of mir-184 mediated angiogenesis are Wnt-receptor Frizzled 7 (Fzd7) and VEGF-A. In our study not only divulges mir-184 as a new and natural inhibitor of LG, but also indicates that multiple mechanisms and factors are involved in maintaining the alymphatic status of the cornea, which warrants further investigation.

Investigative Ophthalmology & Visual Science

Downloaded From: https://arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/934655/ on 12/29/2018
study, we did not detect a significant change in corneal angiogenesis as LG with mir-184 mimic treatment. It is yet to be determined whether factors of the Wnt family and/or VEGF family, such as VEGF-C, are the regulatory components of mir-184 for corneal LG.

Research on mechanisms of corneal LG has broad implications. The use of synthetic microRNA mimic as a treatment, also known as “microRNA replacement therapy,” has emerged as a promising approach for disease treatment and has gained significant attention in cancer therapy. MicroRNA replacement therapy focuses to reintroduce the microRNA to restore loss of function. In this study, using in vivo murine LG model and in vitro human primary LEC culture system, we put forth a similar therapeutic approach for LG interference and suggest mir-184 could be used for the treatment of lymphatic-related diseases, which occur widely inside and outside the eye.

Acknowledgments

Supported in part by research grants from the NIH and University of California at Berkeley (LC).

Disclosure: S. Grimaldo, None; D. Yuen, None; J. Theis, None; M. Ng, None; T. Ecoiffier, None; L. Chen, None

References