Changes in Murine Subbasal Corneal Nerves After Scopolamine-Induced Dry Eye Stress Exposure

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PURPOSE. To evaluate the morphologic changes in the corneal subbasal nerve (CSN) plexus in wild-type (WT) mice after exposure to scopolamine-induced dry eye stress (DES) by using in vivo confocal microscopy.

METHODS. Twenty right eyes of twenty (n = 20) 8-week-old WT BALB/c male mice were investigated. The mice were divided into two experimental groups; 10 eyes of 10 mice exposed to DES for 28 days and 10 eyes of 10 mice were used as a control group. All mice underwent examinations for aqueous tear secretion quantity, tear film breakup time (TBUT), corneal vital staining, and corneal sensitivity thrice (pre-experiment, 2nd week, and 4th week). CSN density, tortuosity, reflectivity, and dendritic cell (DC) densities were examined.

RESULTS. The mean aqueous tear secretion (P < 0.0001) and TBUTs (P < 0.0001) were significantly decreased after DES. The mean corneal vital staining scores were significantly higher (fluorescein, P < 0.0001; lissamine, P < 0.0001), the mean TBUTs were significantly shorter (P < 0.0001), and the corneal sensitivities (P < 0.0001) were significantly lower in the dry eye-induced mice than the control mice. The mean CSN fiber density (P < 0.0001) and the reflectivity (P < 0.0001) were significantly lower; the mean tortuosity and the mean DC density were significantly higher (P < 0.0001) in the dry eye mice.

CONCLUSIONS. Our data demonstrated that prolonged exposure to DES resulted in alterations of CSN density; DC intensity, reflectivity, and tortuosity as well as in tear volume; TBUT; fluorescein and lissamine green staining scores; and the corneal sensitivity in WT mice.

Keywords: corneal subbasal nerves, confocal microscopy, dry eyes, scopolamine

The cornea is an intensively innervated tissue with sensory nerve fiber bundles that maintain corneal epithelial integrity and continuity.1,2 Corneal sensory nerves originate from the ophthalmic branch of the trigeminal ganglion after penetrating the corneal stroma of the corneal limbus. In other words, the cornea is innervated on the intense aggregation of the sensory axon nerve endings called intraepithelial corneal nerves, which one constituted by corneal subbasal nerves (CSNs) and related intraepithelial nerve terminals. The intraepithelial corneal nerve endings have mechanonociceptor, polymodal nociceptor, and cold thermoreceptor functions and stimulate blinking and secreting tears over the cornea.3

Superficially terminating among the corneal epithelial cells, these nerve fibers are easily affected by other ocular or environmental factors.2,4 Subbasal neuronal alterations have been related to various diseases, such as neuropathic corneal ulcers, herpetic keratitis, corneal surgical interventions, diabetes mellitus, and dry eye.5 In such cases, subbasal corneal nerves are damaged with subsequent changes, including reduction of corneal sensitivity, deterioration of corneal epithelial integrity, and decreased in epithelial cell proliferation.5,6

Dry eye disease (DED) has been recently described as a multifactorial chronic disease of the ocular surface characterized by tear film instability, hyperosmolarity, ocular surface inflammation, loss of homeostasis, and neurosensory abnormalities.8 With a better understanding of the complex nature and the significant role of neuropathologic mechanisms in etiology, the association between neurosensory abnormalities and DED was emphasized in the Dry Eye Workshop II (DEWS II) report.9 To evaluate these abnormalities, certain variables, such as CSN density, tortuosity, reflectivity and DCs, have been examined by in vivo confocal microscopy (IVCM).10,11

There are several dry eye mouse models investigating the pathogenesis of DED without considering the subbasal neuronal alterations in dry eye mouse models.12 In our previous experimental study, we generated a short-term dry eye mouse model that mimicked the dry eyes in visual display terminal workers. We called it “the environmental dry eye stress (EDES) model” in which we observed a reduction in CSN density and an increase in dendritic cell (DC) numbers.13 However, there is paucity of knowledge in relation to the effects of longer dry conditions on corneal nerves.

In the current study, we aimed to evaluate the morphologic alterations and branch patterns of the CSN plexus in the wild-type (WT) mice exposed to long-term dry eye stress by IVCM.

MATERIALS AND METHODS

Animals and Micro-Osmotic Pump (MOP) Implantation

Twenty right eyes of 20 (n = 20) 8-week-old WT BALB/c male mice (27–51 g body weight; CLEA, Osaka, Japan) were used for this study. Animals were divided into two experimental groups, namely, 10 eyes of 10 mice exposed to dry eye stress (n = 10).
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The experiment.

repeatability of corneal sensation values was confirmed before

Three measurements were recorded at each time point, and

of both eyes under magnifying glasses without any light source.

mm increments until the mouse demonstrated a complete
corneal blink response. The filament contacted the central

three times by stimulating the corneal nerves to cause a

lowest pressure), each length of nylon filament was applied

to the previous reports.15 Corneal sensitivity assessment was

perpendicularly, avoiding contact with the eyelashes
cornea.

Tokyo, Japan), which had a 0.09-mm diameter, were used to

performed thrice: pre-experiment, 2nd week, and 4th week of

experiencing intervention or often handling of the animals. For subcutane-
ous placement, a minimal incision was performed under the

skin between the muscle tissue. A microcompartment was

created by dissecting the subcutaneous connective tissue apart

with Vannas scissors. The MOP was inserted through the
canthus before repeating the same procedure with the LG dye.

was used to wash out FL, which was removed from the lateral
canthus before repeating the same procedure with the LG dye.

Vital staining photographs were recorded in JPEG format by

using a microscope connected to a digital camera with the

same settings maintained for all mice.

In Vivo Laser Scanning Microscopic Assessment and Analysis of Images

We performed ocular surface confocal analyses by in vivo laser

scanning confocal microscopy with an anterior segment

adaptor. The Rostock Cornea Module of the Heidelberg Retinal

Tomograph II (HRT) (Heidelberg Engineering, Heidelberg,

Germany) was used to examine the corneal morphologic

alterations. Following intraperitoneal anesthesia injection (4

mg/mL of xylazine and 6 mg/mL of ketamine), a carbomer 2%
gel (comfort gel, Dr. Mann Pharma; Fabrik GmbH, Berlin,

Germany) was used to apply the cornea to avoid exposure and
dryness during IVCM examination. The laser source, internally

mounted in the HRT, provides a 670-nm red wavelength diode
di laser. High-resolution real-time images obtained by IVCM

consisted of 384 × 384 pixels covering an area of 400 × 400

µm (horizontal × vertical) with a lateral resolution of 1 µm/
pixel. The images were recorded as a JPEG frame with a

resolution of 8 bits of data and a 128-bit binary floating-point
format. Six to eight complete sequences, each containing 100
images, were recorded from each cornea (each frame representing an area of 160.00 µm²), and the approximate
duration of IVCM examination was 8 minutes per eye. Four

nonoverlapping representative images of each cornea were

selected for corneal morphologic analysis. As previously
described,15 two masked experienced researchers analyzed
the well-focused images for corneal nerve morphologic
features and investigated the density of nerve fibers (NFD),
tortuosity and reflectivity of CSN, and DC density as follows:

1. The NFD was assessed by measuring the total length of
CSN fibers within a frame (160,000 µm²), as defined in
our previous study.15 After semiautomatically marking
the CSN in each frame, the NFD was automatically

Corneal Sensation Assessment

Aqueous Tear Secretion Quantity, Tear Film
Breakup Time (TBUT), and Ocular Surface Vital
Staining Assessments

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for 28 days and 10 eyes of 10 mice used as a control group (n =
10). Sample size calculation for this research study was done
according to the G-Power statistical power analyses program
(Heinrich, Düsseldorf University, Germany) by specifying input
parameters, including the type 1 error, effect size, and standard
deviation.14 To deliver the DES, the mice were shaved from
their back and Alzet MOPs (Model 1002; Durect Corp.,
Cupertino, CA, USA) were implanted subcutaneously just
above their tails for 28 days. The MOP releases prepared
solutions nonstop for 2 weeks without the need for external
intervention or often handling of the animals. For subcutane-
ous placement, a minimal incision was performed under the

skin between the muscle tissue. A microcompartment was

created by dissecting the subcutaneous connective tissue apart

with Vannas scissors. The MOP was inserted through the
incisions that were then sutured with three nonresorbable
sutures. The MOP was replaced with a new one on the 15th
day of the experiment with the same protocol because the
pump life was 14 days. An absolute sterile technique was used
throughout the infilling, interference, and surgical implanta-
tion procedure of the MOP.

Filling MOP With Scopolamine

The MOP has a 0.21 ± 0.01 µL/h mean pumping rate with 100
µL of reservoir volume that can last for 14 days of pumping. To
form DES, 10 mice were infused with 0.2 mg/kg/day scopolamine (dissolved in PBS) per day and 10 mice were
infused with PBS solution as the control group. To avoid
unpredictable pumping rate fluctuations, each pump was filled
completely without trapped air bubbles with a small syringe
(1.0 mL). All experimental procedures were performed under a
fixed room temperature (25 ± 3°C) and humidity (24% ± 4%).
All studies were performed in accordance with the Association
for Research in Vision and Ophthalmology Statement for the
Use of Animals in Ophthalmic and Vision Research. The Animal
Experimentation Ethics Committee of the Keio University
School of Medicine approved the current research procedures.

Aqueous Tear Secretion Quantity, Tear Film
Breakup Time (TBUT), and Ocular Surface Vital
Staining Assessments

All mice underwent examinations for aqueous tear secretion
quantity, TBUT, and corneal vital staining at three time points
(pre-experiment, 2nd week, and 4th week). Aqueous tear secretion quantity, TBUT, and corneal vital staining were
performed according to protocols described in our previous
study.13 Briefly, aqueous tear secretion quantity was evaluated
by phenol red-impregnated cotton threads (Zone-Quick;
Showa Yakuin Kako Co., Ltd., Tokyo, Japan) without
anesthesia. We placed the cotton threads into the lateral
canthus with microforceps for 30 seconds. The wet length of
the cotton threads was measured with a ruler provided by the
manufacturer in millimeters.

TBUT was examined to assess the ocular surface tear film
stability. Initially, one drop of 1 µL of 2% sterile fluorescein (FL)
was applied onto the ocular surface of the mouse by a
micropipette. Excess FL was wiped from the lateral canthus.
After a natural blink response stimulated by air puff by using 1-
µL syringe, the TBUT was examined with portable slit-lamp
biomicroscopy by using cobalt blue light (Kowa Co., Ltd.,
Tokyo, Japan) three times, and the mean of these results was
then calculated.

FL and lissamine green (LG) staining were examined 2
minutes after TBUT assessment by using a scoring system
previously described by Shimamura et al.16 The cornea was
divided into three parts (superior, middle, and inferior), and
each part was scored between 0 and 3 points according to the
extent of the area stained. A score of 0 signified no staining, 1
signified mild staining, 2 signified moderate staining, and 3
signified severe staining for a minimum and maximum total
score of 0 and 9 points, respectively. Five microliters of PBS
was used to wash out FL, which was removed from the lateral
canthus before repeating the same procedure with the LG dye.
Vital staining photographs were recorded in JPEG format by
using a microscope connected to a digital camera with the
same settings maintained for all mice.

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measured by the NeuronJ plug-in for ImageJ software (http://rsb.info.nih.gov/ij/; provided in the public domain by National Institutes of Health, Bethesda, MD, USA). Four different representative images for each cornea were analyzed in pixels. The mean CSN for each cornea was calculated by averaging these total values. The data were determined as density (μm/mm²).

2. The CSN nerve tortuosity was evaluated at the subbasal layer relating to a previous grading scale. The grading scales consisted of a series of images derived from our and other previous studies of corneal innervation using confocal microscopy. Oliveira-Soto et al. 17 classified human corneal nerve tortuosity into five grades. Likewise, in our previous study, we classified mice corneal nerve tortuosity according to subbasal nerve curve magnitude and divided it into five grades from 0 to 5.13

3. The grading of nerve reflectivity was also performed according to our previous research.13 In that study, we classified mice corneal nerve reflectivity into five grades from 0 (no effect) to 5 (strong effect). Similarly, only the nerves longer than 50% of the width frame underwent reflectivity assessment.13

4. The images were also examined for the density of DCs. DCs were morphologically identified through bright cellular images with a branching dendritic morphology located at the level of subbasal nerve plexus layer. Four representative images were used to evaluate the density of epithelial DCs.

Statistical Analysis

To analyze the effects of two factors, including time points and dry eye stress, the 2-way repeated measures ANOVA test was performed to compare tear quantity, vital staining scores, corneal sensation, and CSN parameters at different time points. The Tukey’s test was performed as a multiple comparison test (post hoc test). A P value less than 5% was considered to be statistically significant.

RESULTS

Aqueous Tear Secretion Quantity Alterations

Weight-adjusted aqueous tear production quantity, the mean TBUT, and corneal vital staining score alterations were measured three times during the experiments: before the experiments, 2nd week, and 4th week of the experiments. The mean tear production values for the control group (non-DED) were 0.156 ± 0.05 before the experiment, 0.189 ± 0.04 mm/g at the 2nd week of the experiment, and 0.191 ± 0.04 mm/g at the 4th week of experiment. The mean tear production values for DED group were 0.143 ± 0.02 before the experiment, 0.097 ± 0.01 at 2 weeks of the experiment, and 0.063 ± 0.01 mm/g at 4 weeks of the experiment. Compared with the control group (non-DED), the DED mice demonstrated a significant decrease at the 2nd (P = 0.0081) and 4th (P < 0.0001) week of the experiment in tear volume after exposure to DES (Fig. 1A). At both two and four weeks after exposure to dry eye stress, the mean tear secretion quantity in the DED group was significantly lower than the control group (both time points, P < 0.0001).

TBUT and Ocular Surface Vital Staining Score Alterations

The mean TBUTs for the control group (non-DED) were 2.32 ± 0.40 before the experiment, 2.43 ± 0.30 at the 2nd week of the experiment, and 2.65 ± 0.50 seconds at the 4th week of the experiment. The mean TBUTs in the DED group were
2.05 ± 0.45 before the experiment, 1.30 ± 0.37 at the 2nd week of the experiment, and 0.94 ± 0.29 sec at 4th week of experiment. The mean TBUT in the DED group showed a significant decrease at the 2nd (P = 0.0019) and 4th week (P < 0.0001). (A) Please note a significant increase of FL staining scores in the DED group compared to the vehicle (non-DED) after DES exposure (A, B). (C) The mean FL staining score in the DED group significantly increased after DES exposure at 2nd (P < 0.0001) and 4th week (P < 0.0001). The hash symbol (#) represents comparison between pre-experiment and other time points. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001. The asterisk symbol (*) represents comparison between treatment groups. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001.

Corneal Sensitivity Alterations

Corneal sensitivity in the DES group was significantly lower than the control group at the 2nd week and 4th week of the experiment (2nd week results, 3.5 ± 0.40 in control group (non-DED) versus 2.72 ± 0.30 mm in DED group, P < 0.0001; 4th week results, 3.5 ± 0.30 in control group versus 2.1 ±
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IVCM Results

To investigate whether BALB/c mice also demonstrated any alterations in CSN morphology with DES, we compared CSN density, tortuosity, reflectivity, and DC density between two time points: pretreatment and 4 weeks after exposure to dry eye stress. To assess the effect of DES, a total of 156 images containing 1067 nerves were evaluated for NFD by two experienced researchers (Fig. 4). IVCM assessment for NFD was performed under anesthesia. The mean NFD was 2722 ± 596 in the control (non-DED) group and 2493 ± 208 pixels/frame in the DES group. The data showed that there was no significant change in NFD before the experiment between the two groups (P = 0.38). On the other hand, there was a statistically significant decrease in the mean NFD in the DES group after 4 weeks of the experiment (P = 0.0058). After exposure to 28 days of DES, the mean NFD became 2538 ± 933 in the control (non-DED) group and 1570 ± 506 pixels/frame in the DES group. The mean NFD at 4 weeks after exposure to dry eye stress was significantly lower in the DES group than in the control group (P < 0.0001) (Fig. 5A).

To investigate the tortuosity and reflectivity, 436 nerves from 156 images were investigated using our previously defined grading scale. The mean subbasal nerve tortuosity values were as follows: before experiment control group, 0.52 ± 0.22; before experiment DES group, 0.59 ± 0.15; after 4 weeks of experiment control group, 0.46 ± 0.22; and after 4 weeks of experiment DES group, 1.09 ± 0.25. After exposure to 4 weeks of scopolamine, the mean subbasal nerve reflectivity grades decreased significantly in the DES group (P = 0.011). Moreover, after 4 weeks of experiment there was a statistically significant difference in the mean reflectivity values between the control group and the DES group (P = 0.0028). There was no significant difference in reflectivity values in the control group before and after the experiment (P = 0.765) (Fig. 5C).

The DC density in the control group (8.25 ± 3.59 cells/mm²) was significantly lower than that in the DES group (14.57 ± 4.99 cells/mm²) after 4 weeks of experiment (P = 0.0013). Additionally, there was a significant difference in the DC densities in the DES group before (9.18 ± 4.49) and after the experiment (P = 0.0257) (Fig. 5D).

DISCUSSION

CSNs play an essential role in maintaining ocular surface health and vision as a result of preserving corneal integrity and sensation. Recently, IVCM has become a widely used noninvasive diagnostic tool for the clinical evaluation of the cornea, which ensures high-resolution images of corneal nerves, structures, and layers. Jiang et al. reported that noninvasive IVCM imaging of CSN is a practical and useful method to determine the alterations of small fiber neuropathy. Because previous studies have reported a high agreement in the evaluation of corneal nerves between IVCM and histologic examinations, IVCM has been used to evaluate qualitative and quantitative variations of the corneal nerves in many conditions, including herpes zoster ophthalmicus, corneal transplantation, laser refractive surgery (laser-assisted epithelial keratomileusis and photorefractive keratectomy), noncorneal ocular procedures (cataract surgery, laser panretinal photocoagulation, laser retinopexy, and scleral buckles), antiglaucoma medical therapy (particularly topical beta-adrenergic antagonists), aging, diabetes mellitus, and dry eye. A significant decrease in CSN density has been shown nearly in all of these conditions. Most of the studies investigating corneal nerves in DED patients also mainly assessed the CSN density and found a decrease in the majority, whereas others found an increase or no change. These variable results were found to be associated with differences in DED severity, corneal nerve damage severity, neural regeneration/degenerative patterns,
and inflammation levels. Recently, Stepp et al. have demonstrated that decreased intraepithelial corneal nerves in the CD25 null Sjögren syndrome mouse model is accompanied by reduced corneal sensitivity and increased corneal epithelial cell proliferation. Furthermore, Stepp et al. reported a significant decrease in axon thickness, density, and apical extension of the intraepithelial nerve terminals after 5 days of dry eye exposure in C57BL/6 mice. Additionally, a previous
study done by De Paiva et al. has shown acute deterioration of barrier function and increased inflammation even after 5 days of desiccating stress in mice. Similarly, we found a notable decrease in subbasal nerve density after acute environmental stress in our previous study. In the current study, 28 days of dry eye stress reduced CSN density in the WT mice.

In previous dry eye studies, besides CSN density, other morphologic parameters, including tortuosity, reflectivity, DC density, and beading pattern, have been also been evaluated with variable results (related to neural regenerations). Most of these human studies performed in patients with dry eye have shown an increase in DC density and alterations in other parameters due to the consequence of neural regeneration and inflammation stimulated by both innate and adaptive immune responses is critical to pathogenesis and the chronicity of DED. In the current study, we also found an increase in nerve tortuosity and a reduction in nerve reflectivity in the dry eye patients. An increase in tortuosity in chronic stress was considered to be associated with neural regenerations, as demonstrated in previous studies on chronic diseases, such as diabetic neuropathy, keratoconus, Sjögren’s syndrome, DED, and meibomian gland dysfunction, and our results were consistent with these studies.

Moreover, in this study, we also evaluated tear volume, TBUT, FL staining scores, LG staining scores, and corneal sensitivity changes after DES exposure. Both acute and longer term dry eye stress showed a similar level of decreased tear secretion quantity. We observed a significant increase in FL and LG staining scores, and corneal sensitivity changes after DES exposure. The hash symbol (#) represents comparison between time points. *P < 0.05, **P < 0.01, ###P < 0.001, and ####P < 0.0001. The asterisk symbol (*) represents comparison between treatment groups.

As for tortuosity and reflectivity, the results were different between acute and longer term stress conditions. In this study, we found an increase in tortuosity and decrease in reflectivity after exposure to dry eye stress, whereas these parameters were stable in our previous acute EDES study. Recently, a dry eye study comparing CSN tortuosity and reflectivity between dry eye patients and controls also has found an increase in tortuosity and a reduction in nerve reflectivity in the dry eye patients. An increase in tortuosity in chronic stress was considered to be associated with neural regenerations, as demonstrated in previous studies on chronic diseases, such as diabetic neuropathy, keratoconus, Sjögren’s syndrome, DED, and meibomian gland dysfunction, and our results were consistent with these studies.

This study has several limitations, including the use of only male mice (to avoid hormonal changes of menstrual cycle). Furthermore, we did not observe how the nerves healed after longer term DES. Further studies should be performed to evaluate the chronic effect of DES. In this study, we used the BALB/c mice strain because CSN density is higher in BALB/c mice compared with same-aged C57BL/6 mice. Further studies should also focus on female mice and different mouse strains. In this study, we evaluated the alterations of DC
density. Although DCs were carefully identified by two examiners through bright cellular images with branching dendritic morphology, they were not phenotypically determined as DCs. Further study using immunohistochemistry should be performed to confirm our current results.

In the human study, it is difficult to examine the sole effect of dry eye on the corneal nerve because the influence of the systemic diseases and external factors cannot be excluded. Therefore, there are discrepancies in reporting nerve alterations between DE studies as described above. In this respect, conducting research by using a dry eye animal model is essential for understanding the pathophysiology. The novelty of this study is that we observed in vivo alterations in the subepithelial nerve plexus after scopolamine-induced dry eye stress in a mouse model. This study may form the basis of future research to evaluate the dry eye treatment drugs and elucidate the pathophysiology of the disease.

In conclusion, prolonged exposure to DES in WT mice resulted in significant changes in CSN density, DC intensity, reflectivity, and tortuosity as well as in tear volume TBUFL, FL-LG staining scores, and corneal sensitivity.

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